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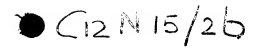
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- (S) Use of oxidoreductases in bleaching and/or detergent compositions and their preparation by microorganisms engineered by recombinant DNA technology.
- The structural genes and their regulatory DNA sequences of an alcohol oxidase (MOX) and a dihydroxyacetone synthase (DHAS) of Hansenula polymorpha have been isolated and the nucleotide sequences determined. The invention relates to the use of the MOX gene, as well as the use of the regulatory DNA sequences of MOX and/or DAS in combination with the MOX gene, optionally after modification thereof, or other oxidase genes, or other genes, to produce engineered microorganisms, in particular yeasts.

Said engineered microorganisms can produce oxidases or other enzymes in yields that allow industrial application on a large scale.

Moreover, said engineered microorganisms can produce oxidases having improved properties with respect to their application in oxidation reactions and/or in bleaching and detergent products.

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USE OF OXIDOREDUCTASES IN BLEACHING AND/OR DETERGENT COMPOSITIONS AND THEIR PREPARATION BY MICROORGANISMS ENGINEERED BY RECOMBINANT DNA TECHNOLOGY

The present invention relates to a process for microbiologically preparing oxidoreductases, use of these enzymes in bleaching and/or detergent compositions, as well as to microorganisms transformed by DNA sequences coding for an oxidoreductase and optionally for a dihydroxyacetone synthase-enzyme, and H. polymorpha alcohol oxidase and/or dihydroxyacetone synthase regulation sequences, the microorganisms being suitable for use in the process.

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Oxidoreductases, especially those which use oxygen as electron acceptor, are enzymes suitable for use in bleaching and/or detergent compositions in which they can be used for the <u>in situ</u> formation of bleaching agents, e.g.  $\rm H_2O_2$ , during the washing or bleaching process. See for example

- GB-PS 1 225 713 (Colgate-Palmolive Company), in which the use of a mixture of glucose and glucose oxidase and other ingredients in a dry powdered detergent composition has been described,
- DE-PA 2 557 623 (Henkel & Cie GmbH), in which the use of a C<sub>1</sub> to C<sub>3</sub> alkanol and alcohol oxidase, or galactose and galactose-oxidase, or uric acid and uratoxidase, and other ingredients in a dry detergent composition having bleaching properties has been described, and
- GB-PA 2 101 167 (Unilever PLC) in which the use of a  $C_1$  to  $C_4$  alkanol and a  $C_1$  to  $C_4$  alkanol oxidase in a liquid bleach and/or detergent composition has been described,

wherein the alkanol and the enzyme are incapable of substantial interaction until the composition is diluted with water, and/or has come into contact with sufficient oxygen.

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Up to now natural oxidase-enzymes cannot be produced at a cost price that allows industrial application on a large scale, e.g. detergent products. Moreover, the oxidase-enzymes have to act under non-physiological conditions when used in detergent and bleaching products. Further the natural oxidases that have been investigated for use in detergent compositions are accompanied by the natural catalase-enzyme which decomposes almost immediately the peroxide(s) formed, so that no effective bleaching is obtained. Thus a need exists for oxidase-enzymes that are more suitable for use under the conditions of manufacture and use of detergent and bleaching products.

For an economically feasible production of these oxidases it is further required to reach a yield of these enzymes in fermentation processes in the order of that of alcohol oxidase of <u>H. polymorpha</u>, which is up to 20% of the cellular protein (van Dijken et al., 1976).

One way of finding new microorganisms producing enzymes in higher amounts or finding new oxidase-enzymes having improved properties is to check all sorts of microorganisms and try to isolate the relevant oxidases, which are then checked for their abilities to generate peroxides and their stabilities under the conditions of manufacture and use of detergent and bleaching products. One can hope that some day a suitable enzyme will be found, but the chance of success is unpredictable and probably very low.

Another way is to apply another trial and error method of crossing the natural microorganisms producing these oxidases by classical genetic techniques, in the hope that some day one will find a more productive microorganism or a more suitable enzyme, but again the chance of success is rather low.

Clearly, a need exists for a method for preparing 5 oxidase-enzymes in higher yield and/or without the concomitant formation of catalase and/or having improved properties during storage and/or use in e.g. bleach and/or detergent compositions. The problem of trial and error can be overcome by a process for preparing an 10 oxidase-enzyme by culturing a microorganism under suitable conditions, and preferably concentrating the enzyme and collecting the concentrated enzyme in a manner known per se, which process is characterized in that a microorganism is used that has been obtained by 15 recombinant DNA technology and which is capable of producing said oxidase-enzyme.

The microorganisms suitable for use in a process for preparing an oxidase-enzyme can be obtained by recombinant DNA technology, whereby a microorganism is transformed by a DNA sequence coding for an oxidase-enzyme (so-called structural gene) together with one or more other DNA sequences which regulate the expression of the structural gene in a particular microorganism or group of organisms, either via introduction of an episomal vector containing said sequences or via a vector containing said sequences which is also equipped with DNA sequences capable of being integrated into the chromosome of the microorganism.

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The determination of a structural gene coding for the enzyme alcohol oxidase (EC 1.1.3.13) originating from H. polymorpha together with its regulatory 5'- and 3'-flanking regions will be described as an example of the invention without the scope of the invention being limited to this example. The spirit of the invention is

also applicable to the isolation of DNA sequences of other oxidase-enzymes such as glycerol oxidase, glucose oxidase, D-amino acid oxidase etc.; the incorporation of the DNA sequences or modifications thereof into the genome of microorganisms or into episomal vectors used for transforming microorganisms and the culturing of the transformed microorganisms so obtained as such or for producing the desired oxidase-enzymes, as well as the use of these enzymes in bleaching compositions containing them.

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Although the microorganisms to be used can be bacteria, e.g. of the genus Bacillus, as well as moulds, the use of yeasts is preferred for technological and economical reasons. In particular a mould or yeast can be selected from the genera Aspergillus, Candida, Geotrichum, Hansenula, Lenzites, Nadsonia, Pichia, Poria, Polyporus, Saccharomyces, Sporobolomyces, Torulopsis, Trichosporon and Zendera, more particularly from the species A. japonicus, A. niger, A. oryzae, C. boidinii, H. polymorpha, Pichia pastoris and Kloeckera sp. 2201. The latter name is sometimes used instead of C. boidinii.

Many C<sub>1</sub>-utilizing yeasts have been isolated during the last decade, and for <u>Hansenula polymorpha</u> and <u>Candida boidinii</u> the methanol metabolism has been studied extensively (for a review see Veenhuis et al., 1983).

The first step in this metabolism is the oxidation of methanol to formaldehyde and  ${\rm H_2O_2}$  catalysed by MOX. Formaldehyde is oxidized further by the action of formaldehyde dehydrogenase and formate dehydrogenase.

 $H_2O_2$  is split into water and oxygen by catalase.

Alternatively, methanol is assimilated into cellular

material. After its conversion into formaldehyde, this product is fixed through the xylulose monophosphate pathway into carbohydrates. Dihydroxyacetone synthase (DHAS) plays a crucial role in this assimilation process.

The appearance of MOX, formate dehydrogenase, formal-dehyde dehydrogenase, DHAS and catalase is subject to glucose repression, e.g. on 0.5% glucose. However, synthesis of MOX is derepressed by growth in low concentrations of glucose (0.1%), contrary to the synthesis of DHAS, which is still fully repressed under these conditions (Roggenkamp et al., 1984).

Regulation, i.e. the possibility to switch "on" or "off" of the gene for the polypeptide concerned, is desirable, because it allows for biomass production, when desired, by selecting a suitable substrate, such as, for example melasse, and for production of the polypeptide concerned, when desired, by using methanol or mixtures of methanol and other carbon sources.

Methanol is a rather cheap substrate, so the polypeptide production may be carried out in a very economical way.

After derepression of the gene coding for alcohol oxidase (MOX) by growth on methanol, large microbodies, the peroxisomes are formed. While glucose-grown cells contain only a small peroxisome, up to 80% of the internal volume of the cell is replaced by peroxisomes in the derepressed state. The conversion of methanol into formaldehyde and  $\rm H_2O_2$  as well as the degradation of  $\rm H_2O_2$  has been shown to occur in these peroxisomes, while further oxidation or assimilation of formaldehyde most probably occurs in the cytoplasm. This process is a perfect example of compartmentalization of toxic pro-

ducts, of a strong co-ordinate derepression of several cellular processes and of the selective translocation of at least two of the enzymes involved in this process.

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Most of the enzymes involved in the methanol metabolism have been purified and characterized (Sahm, 1977, Bystrykh et al, 1981). Especially methanol oxidase (EC 1.1.3.13) has been studied in detail. It is an octamer consisting of identical monomers with an Mr value of about 74 kd and it contains FAD as a prosthetic group. Up to now no cleavable signal sequence for translocation could be detected, as concluded from electroelephoresis studies with in vivo and in vitro synthesized products (Roa and Blobel, 1983) or from in vitro synthesis in the presence of microsomal membranes (Roggenkamp et al., 1984).

Under derepressed conditions, up to 20% of the cellular 20 protein consists of MOX.

### Materials and methods

- A) Microorganisms and cultivation conditions

  Hansenula polymorpha CBS 4732 was obtained from Dr

  J.P. van Dijken (University of Technology, Delft,
  The Netherlands). Cells were grown at 37°C in 1
  litre Erlenmeyer flasks containing 300 ml minimal
  medium (Veenhuis et al., 1978), supplemented with

  0.5% (v/v) methanol or 0.5% (v/v) ethanol as
  indicated. Phage lambda L47.1 and the P2 lysogenic

  E. coli K12 strain Q 364 were obtained from Dr P.
  van der Elsen (Free University of Amsterdam, The
  Netherlands) and propagated as described (Loenen and
  Brammar, 1980).
  - E. coli K12 strains BHB 2600, BHB 2688 and BHB 2690

(Hohn, 1979) were obtained from Dr M. van Montagu (University of Gent, Belgium), while <u>E. coli</u> Kl2 strain JM 101.7118 and the Ml3 derivatives Ml3 mp 8, 9, 18 and 19 were obtained from Bethesda Research Laboratories Inc. (Gaithersburg, MD, U.S.A.).

#### b) Enzymes '

All enzymes used were obtained from Amersham International PLC, Amersham, U.K., except alpha-helicase which was obtained from Pharm Industrie, Clichy, France. Enzyme incubations were performed according to the instructions of the manufacturer.

ATP:RNA adenyl transferase was purified as described by Edens et al. (1982).

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C) Other materials
[35s] methionine, [alpha-35s] dATP, [alpha-32p]
dNTP's, [alpha-32p] ATP and [gamma-32p] ATP were
obtained from Amersham International PLC, Amersham,
U.K.

Nitrobenzyloxy-methyl (NBM) paper was obtained from Schleicher and Schuell, and converted into the diazo form (DBM) according to the instructions of the manufacturer.

Nitrocellulose filters (type HATF) were obtained from Millipore.

# 30 RNA isolation, fractionation and analysis

Hansenula polymorpha cells were grown to midexponential phase, either in the presence of methanol
or ethanol. The cells were disrupted by forcing them
repeatedly through a French Press at 16 000 psi, in a
buffer containing 10 mM Tris-HCl pH 8, 5 mM MgCl<sub>2</sub>,
1% NaCl, 6% para-aminosalicylic acid, 1% sodium do-

decylsulphate (SDS) and 5% phenol. The purification of polyadenylated RNA was subsequently performed, as described previously (Edens et al., 1982). One gram cells yielded four mg total RNA and 0.1 mg polyadenylated RNA. Five microgram samples of total RNA or polyadenylated 5 RNA were radioactively labelled at their 3'-ends with ATP: RNA adenyl transferase and [alpha-32p] ATP, and subsequently separated on a 2.5% polyacrylamide gel containing 7 M urea (Edens et al., 1982). For the preparative isolation of a specific mRNA fraction, 40 10 micrograms polyadenylated RNA was mixed with four micrograms of labelled polyadenylated RNA and separated on the denaturing polyacrylamide gel. The radioactive 2.4 kb RNA class was eluted from slices of the gel and freed from impurities by centrifugation through a 5-30% 15 glycerol gradient in 100 mM NaCl, 10 mM Tris-HCl pH 7.5, 1 mM EDTA and 0.1% SDS for 15 h at 24 000 rev./min. in a Beckmann centrifuge using an SW 60 rotor at 20°C. The radioactive fractions were pooled and 20 precipitated with ethanol. Polyadenylated RNA was translated in vitro in a rabbit reticulocyte lysate according to Pelham and Jackson (1976), using [358] methionine as a precursor. The translation products were immuno-precipitated with MOX antiserum as des-25 cribed by Valerio et al. (1983).

### cDNA synthesis

One third of the RNA fraction, isolated from the polyacrylamide gel, was used to procure a radioactive cDNA
with reverse transcriptase (Edens et al., 1982). Using
[alpha-32P] dATP and [alpha-32P] dCTP of a high
specific activity (more than 3000 Ci/mM), 20 000 cpm of
high molecular weight cDNA was formed during 1 h at
42°C in the presence of human placental ribonuclease
inhibitor.

#### DNA isolation

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Ten g of Hansenula polymorpha cells were washed with 1 M sorbitol and resuspended in 100 ml 1.2 M sorbitol, 10 mM EDTA and 100 mM citric acid pH 5.8, to which 100 5 microliter beta-mercapto-ethanol was added. Cells were spheroplasted by incubation with 500 mg alpha-helicase for 1 h at 30°C. Spheroplasts were collected by centrifugation at 4000 rev./min. in a Sorvall GSA rotor, resuspended in 40 ml 20 mM Tris-HCl pH 8, 50 mM EDTA 10 and lysed by adding 2.5% SDS. Incompletely lysed cells were pelleted for 30 min. at 20 000 rev./min. in a Sorvall SS34 rotor and DNA was isolated from the viscous supernatant by centrifugation using a CsClethidium bromide density gradient at 35 000 rev./min. 15 for 48 h in a Beckmann centrifuge using a 60 Ti rotor. 2 mg of DNA was isolated with a mean length of 30 kb.

# Preparation of a clone bank in phage lambda L47.1

150 microgram Hansenula polymorpha DNA was partially digested with Sau3AI and sedimented through a 10-40% sucrose gradient in 1 M NaCl, 20 mM Tris-HCl pH 8 and 5 mM EDTA for 22 h at 23 000 rev./min. in an SW 25 rotor. The gradient was fractionated and samples of the fractions were separated on a 0.6% agarose gel in TBE buffer (89 mM Tris, 89 mM Boric acid, 2.5 mM EDTA).

Fractions that contained DNA of 5-20 kb were pooled and the DNA was precipitated with ethanol. Phage lambda L47.1 was grown, and its DNA was isolated as described by Ledeboer et al. (1984). The DNA was digested with BamHI and arms were isolated by centrifugation through a potassium acetate gradient as described by Maniatis et al. (1982). Two microgram phage lambda DNA arms and 0.5 /ug Sau3AI digested Hansenula polymorpha DNA thus obtained were ligated and packaged in vitro using a

protocol from Hohn (1979). Phages were plated on  $\underline{E}$ .  $\underline{\text{coli}}$  strain Q 364 to a plaque density of 20,000 pfu per 14 cm Petri dish. Plaques were blotted onto a nitrocellulose filter (Benton and Davis, 1977) and the blot was hybridized with the radioactive cDNA probe isolated as described above. Hybridization conditions were the same as described by Ledeboer et al. (1984) and hybridizing plaques were detected by autoradiography.

# 10 <u>Isolation and partial amino acid sequence analysis of alcohol oxidase (MOX)</u>

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Hansenula polymorpha cells grown on methanol were disintegrated by ultrasonification and the cell debris was removed by centrifugation. The MOX-containing protein fraction was isolated by  $(NH_4)_2SO_4$  precipitation . (40-60% saturation). After dialysis of the precipitate, MOX was separated from catalase and other proteins by ion-exchange chromatography (DEAE-Sepharose) and gel filtration (Sephacryl S-400). Antibodies against MOX were raised in rabbits by conventional methods using complete and incomplete Freund's adjuvants (Difco Lab, Detroit, U.S.A.). Sequence analysis of alcohol oxidase treated with performic acid was performed on a Beckman sequenator. Identification of the residues was done with HPLC. The amino acid composition was determined on a Chromaspek analyser (Rank Hilger, U.K.), using standard procedures and staining by ninhydrine. The carboxy terminal amino acid was determined as described by Ambler (1972).

# Chemical synthesis of deoxyoligonucleotides

Deoxyoligonucleotides were synthesized on a Biosearch

SAM I gene machine, using the phosphite technique

(Matteucci and Caruthers, 1981). They were purified on

16% or 20% polyacrylamide gels in TBE.

# Hybridization with deoxyoligonucleotide probes

The deoxyoligonucleotides were radioactively labelled with  $T_4$ -polynucleotide kinase and [gamma- $^{32}$ P] ATP. The DNA of the MOX clones obtained was digested with different restriction enzymes, separated on 1% agarose gel and blotted onto DBM paper. Hybridizations were performed as described by Wallace et al. (1981).

### 10 DNA sequence analysis

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From clone 4 (see Example 1) containing the complete MOX gene, several subclones were made in phage Ml3mp-8, -9 or M13mp-18, -19 derivatives by standard techniques. Small subclones (less than 0.5 kb), cloned in two 15 orientations, were sequenced directly from both sides. From the larger subclones, also cloned in two orientations, sequence data were obtained by an exonuclease Bal31 digestion strategy (see Fig. 1). For each of both cloned orientations the RF Ml3 DNA is digested with a 20 restriction enzyme that preferably cleaves only in the middle of the insert. Subsequently, both orientations of the clones were cut at this unique site, and digested with exonuclease Bal31 at different time intervals. Incubation times and conditions were chosen such that 25 about 100-150 nucleotides were eliminated during each time interval. Each fraction was digested subsequently with the restriction enzyme, recognizing the restriction site situated near the position at which the sequence reaction is primed in the Ml3 derivatives. Ends were made 30 blunt end by incubation with  $T_A$ -polymerase and all dNTP's, and the whole mix was ligated under diluted conditions, thereby favouring the formation of internal RF molecules. The whole ligation mix was used to transform to E. coli strain JM 101-7118. From each time 35 interval several plaques were picked up and sequenced using recently described modifications of the Sanger sequencing protocol (Biggin et al., 1983).

# The isolation of auxotrophic mutants

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LEU-1 (CBS N° 7171) is an auxotrophic derivative of H. polymorpha strain NCYC 495 lacking  $\beta$ -isopropylmalate dehydrogenase activity. The isolation of this mutant has been described by Gleeson et al. (1984).

LR9 (CBS N° 7172) is an auxotrophic derivative of H. polymorpha ATCC 34438, lacking orotidine 5'-decarbox-ylase activity.

For the isolation, all procedures were carried out at 30°C instead of 37°C, which is the optimal temperature for growth of this yeast. Yeast cells were mutagenized 15 with 3% ethylmethanesulphonate for 2 hr (Fink, 1970). The reaction was stopped with 6% sodium thiosulphate (final concentration) and the solution was incubated for another 10 min. Mutagenized cells were then washed once with H2O and incubated for 2 days on YEPD or YNB supple-20 mented with uracil for segregation and enrichment of uracil-auxotrophs followed by a 15 hr cultivation on MM without nitrogen source. Finally a nystatin enrichment was employed for 12 hr on MM with a concentration of 10 jug antibiotic per ml. The treated cells were plated 25 on YNB plates containing 200 jug uracil per ml and 0.8 mg 5-fluoroorotic acid (Boeke et al., 1984). Usually 10<sup>6</sup> cells were plated on a single plate. Resistant colonies were picked after 3 days of incubation, replica plated twice on YNB plates to establish the auxotrophy. From the auxotrophic mutants ura 30 cells were isolated. Alternatively, 1.5 x 10<sup>6</sup> yeast cells were incubated in one ml of YNB liquid medium supplemented with 200 jug of uracil and 0.8 mg of 5fluoroorotic acid. After incubation of 2 days, the treated cells were plated on YNB containing uracil, 35 replica-plated twice on YNB and analysed as described above.

Such resistant mutants have been shown to be uracil auxotrophs affected at the URA3 or the URA5 locus in  $\underline{S}$ . cerevisiae (F. Lacroute, personal communication). Of about 600 resistant colonies of H. polymorpha tested, 52 exhibited a uracil phenotype. Since URA3 and URA5 5 mutations in S. cerevisiae lack orotidine 5'decarboxylase and orotidine 5'-phosphate pyrophosphorylase, respectively (Jones and Fink, 1982), the obtained uracil auxotrophs of H. polymorpha were tested for both enzymatic activities (Lieberman et al., 10 1955). Mutants affected in either of the two enzymes were found (Table I). They have been designated odcl and oppl mutants, respectively. The odcl mutants exhibit adequate low reversion frequencies (Table II) and thus are suitable for transformation purposes by com-15 plementation.

# Isolation of autonomous replication sequences (HARS) from H. polymorpha

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Chromosomal DNA from <u>H</u>. <u>polymorpha</u> was partially digested either with <u>Sal</u>I or <u>Bam</u>HI and ligated into the single <u>Sal</u>I and <u>Bam</u>HI site of the integrative plasmid YIp5, respectively. The ligation mixture was used to transform <u>E</u>. <u>coli</u> 490 to ampicillin resistance. YIp5 is an integrative plasmid containing the URA3 gene as a selective marker (Stinchcomb et al., 1980).

The plasmid pool of  $\underline{H}$ . polymorpha SalI clones was used to transform  $\underline{H}$ . polymorpha mutant LR9. A total of 27 transformants was obtained being also positive in the  $\beta$ -lactamase assay. From all of them, plasmids could be recovered after transformation of  $\underline{E}$ . coli 490 with yeast minilysates. Restriction analysis of the plasmids revealed that most of the inserts show the same pattern. The two different plasmids, pHARS1 and pHARS2, containing inserts of 0.4 and 1.6 kb respectively, were

used for further studies (Fig. 2). Both plasmids transform H. polymorpha mutant LR9 with a frequency of about 500-1,500 transformants per /ug of DNA using the transformation procedure of intact cells treated with polyethyleneglycol. Southern analysis of the H. polymorpha transformants after retransformation with pHARS1 and pHARS2 recovered from E. coli plasmid preparations shows the expected plasmid bands and thus excludes integration of the URA3 gene as a cause of the uracil protrophy. Therefore, we conclude that the HARS sequences like ARS1 (Stinchcomb et al., 1982) allow autonomous replication in H. polymorpha. Neither HARS1 nor HARS2 enabled autonomous replication in S. cerevisiae. HARS1 was sequenced completely as shown in Fig. 3.

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# Estimation of plasmid copy number in H. polymorpha transformants

The copy number of plasmids conferring autonomous rep-20 lication in H. polymorpha either by ARS sequences or by HARS sequences was estimated by Southern blot analysis (Fig. 4). For comparison, plasmid YRP17 in S. cerevisiae (Fig. 4, lanes 6, 7), which has a copy number 25 of 5-10 per cell (Struhl et al., 1979) and the high copy number plasmid pRB58 in S. cerevisiae (Fig. 4, lanes 4, 5) with about 30-50 copies per cell were used. YRP17 is a URA3-containing yeast plasmid, bearing an ARS sequence (Stinchcomb et al., 1982), while pRB58 is a 2 /um derivative containing the URA3 30 gene (Carlson and Botstein, 1982). A Kluyveromyces lactis transformant carrying 2 integrated copies of pBR pBR322 was used as a control (Fig. 4, lanes 2, 3). The intensity of staining in the autoradiogram reveals that the plasmid YRP17 in H. polymorpha has practically 35 the same copy number as in S. cerevisiae, whereas plasmids pHARS-1 and pHARS-2 show a copy number which is in

the range of about 30-40 copies per cell like pBR58 in S. cerevisiae. This proves once more the autonomously replicating character of the HARS sequence.

# 5 Transformation procedures

Several protocols were used.

a) H. polymorpha strain LEU-1 was transformed using a procedure adapted from Beggs (1978). The strain was 10 grown at 37°C with vigorous aeration in 500 ml YEPD liquid medium up to an OD<sub>600</sub> of 0.5. The cells were harvested, washed with 20 ml distilled water and resuspended in 20 ml 1.2 M sorbitol, 25 mM EDTA pH 8.0, 150 mM DTT and incubated at room tem-15 perature for 15 minutes. Cells were collected by centrifugation and taken up in 20 ml 1.2 M sorbitol, 0.01 M EDTA, 0.1 M sodium citrate pH 5.8 and 2% v/vbeta-glucuronidase solution (Sigma 1500000 units/ml) and incubated at 37°C for 105 minutes. After 1 hr, 20 the final concentration of beta-glucuronidase was brought to 4% v/v. For transformation, 3 ml aliquots of the protoplasts were added to 7 ml of ice cold 1.2 M sorbitol, 10 mM Tris-HCl pH 7. Protoplasts were harvested by centrifugation at 2000 rpm for 5 25 minutes and washed three times in ice cold sorbitol buffer. Washed cells were resuspended in 0.2 ml 1.2 M sorbitol, 10 mM  $\operatorname{CaCl}_2$ , 10 mM  $\operatorname{Tris-HCl}$  pH 7 on ice. 2 /ug of YEP13 DNA - an autonomous replicating S. cerevisiae plasmid consisting of the LEU2 30 gene of S. cerevisiae and the 2 micron-ori (Broach et al., 1979) - were added to 100 ml of cells and incubated at room temperature. 0.5 ml of a solution of 20% PEG 4000 in 10 mM CaCl2, 10 mM Tris-HCl pH 7.5 was added and the whole mixture was incubated 35 for 2 minutes at room temperature. Cells were collected by brief (5 sec.) centrifugation in an MSG

microfuge set at high speed and resuspended in 0.1 ml YEPD 1.2 M sorbitol pH 7.0, and incubated for 15 minutes at room temperature. The cells were plated directly by surface spreading on plates containing 2% Difco agar, 2% glucose, 0.67% Difco yeast nitrogen base and 20 mg/l of each of L-adenine Hemisulphate, methionine, uracil, histidine, tryptophan, lysine and 1.2 M sorbitol. Leu<sup>+</sup> transformants appear after 5 days incubation at 37°C with a frequency of 50 colonies/ug DNA, while no transformants appear if no DNA is added.

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- b) Alternatively, H. polymorpha LEU-1 was transformed with YEP13, using a procedure adapted from Das et al. (1984). Exponentially growing cells were grown 15 up to an OD<sub>600</sub> of 0.4, washed in TE buffer (50 mM Tris-HCl pH 8.0, 1 mM EDTA) and resuspended in 20 ml TE buffer. 0.5 ml cells were incubated with 0.5 ml 0.2 M LiCl for 1 hr at 30°C. To 100 ml of these 20 cells 4 /ug YEP13 in 20 ml TE buffer was added and the sample was incubated for a further 30 minutes at 30°C. An equal volume of 70% v/v PEG 4000 was added and the mixture was incubated for 1 hr at 30°C, followed by 5 min. at 42°C. After addition of 1 ml 25 H<sub>2</sub>O, cells were collected by a brief centrifugation as described under a), washed twice with H<sub>2</sub>O and resuspended in 0.1 ml YEPD 1.2 M sorbitol and incubated for 15 minutes at room temperature. Cells were plated as described. Leu+ transformants 30 appear with a frequency of 30/jug DNA.
- c) The H. polymorpha URA mutant LR9 was transformed with YRP17, a plasmid containing the URA3 gene of S. cerevisiae as a selective marker and an autonomously replicating sequence (ARS) for S. cerevisiae (Stinchomb et al, 1982). Using the protoplast method described by Beggs (1978), 2-5 transformants/jug

DNA were obtained. This number was enlarged, using the LiSO<sub>4</sub> method of Ito et al. (1983), up to 15-20 transformants per /ug of DNA. However, the best procedure was the procedure described by Klebe et al. (1983), using intact cells treated with PEG 4000. Up to 300 transformants were obtained per /ug DNA. The LiSO<sub>4</sub> procedure, as well as the Klebe procedure, was performed at 37°C.

Transformation of <u>H. polymorpha</u> based on autonomous replication of the vector was indicated by two characteristics: (1) the instability of the uracil<sup>+</sup> phenotype. After growth of transformants on YEPD for ten generations, more than 99% had lost the ability to grow on selective medium (Table II). (2) Autonomous replication was further ascertained by transforming <u>E. colicells</u> with yeast minilysates and retransformation of <u>H. polymorpha</u>. Subsequent Southern analysis showed the presence of the expected plasmid.

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H. polymorpha LR9 could not be transformed with pRB58, or with pHH85, constructed by insertion of the whole 2 micron circle DNA (Hollenberg, 1982) into the PstI site of the ampicillin gene of plasmid YIP5. YIP5, containing the DNA sequence of HARS1 or HARS2, was transferred to H. polymorpha LR9 using the Klebe protocol with a frequency of 500-1500 transformants per /ug of DNA. Thus, transformation frequency is 2-5 times higher than described above, using the heterologous ARS 1 in YRP17 of S. cerevisiae. Similarly, the stability of the HARS plasmid in transformants is slightly higher than the ARS 1 plasmid (Table II).

Transformation of H. polymorpha by integration of the URA3 gene from S. cerevisiae

The URA3 gene of  $\underline{S}$ . cerevisiae shows no homology to the

ODC gene in H. polymorpha, as revealed by Southern hybridisation of nick-translated YIP5 plasmid DNA to chromosomal DNA of H. polymorpha. Therefore, low-frequency integration of the URA3 gene at random sites of the H. polymorpha genome had to be anticipated. Transformation of mutant LR9 with the integrative vector YIP5 resulted in 30-40 colonies per /ug of DNA on YNB plates using the polyethyleneglycol method, whereas no transformants were obtained in the control experiment using YIP5 for transformation of S. cerevisiae mutant YNN27. Analysis of 38 transformants revealed 4 stable integrants after growth on non-selective medium. The integration event was further demonstrated by Southern analysis (Fig. 5).

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A second procedure for generating integration of the URA3 gene into chromosomal DNA of H. polymorpha was performed by enrichment of stable Ura+ transformants from transformants carrying plasmid pHARS1. Transformants were grown in liquid YEPD up to a density of  $10^9$  cells per ml. An aliquot containing 5 x  $10^6$ cells was used to inoculate 100 ml of fresh medium and was grown up to a cell density of 109 per ml. The procedure was repeated until about 100 generations had been reached. Since the reversion rate of mutant LR9 is  $2 \times 10^{-9}$  and the frequency of plasmid loss per 10 generations is 97% in pHARS1 transformants, the predominant part of the Ura + cells after 100 generations should be integrants. The Ura colonies tested were all shown to maintain a stable Ura phenotype indicating an integration of the URA3 gene. This was further verified by Southern blot analysis. In addition, these data indicate that the integration frequency is  $5 \times 10^{-6}$ .

### Example 1

CLONING OF THE GENE FOR ALCOHOL OXIDASE (MOX) FROM HANSENULA POLYMORPHA

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# Characterization of polyadenylated RNA

Total RNA and polyadenylated RNA, isolated from cells grown on methanol, were labelled at their 3'-termini with 10 ATP: RNA adenyl transferase, and separated on a denaturing polyacrylamide gel (Fig. 6). Apart from the rRNA bands, two classes of RNA appear in the poly-adenylated RNA lane, respectively 1 kb and 2.3 kb in length. Since these RNA classes are not found in polyadenylated RNA of 15 ethanol-grown cells (result not shown), they obviously are transcripts of genes derepressed by growth on methanol. The 2.3 kb class can code for a protein of 700 to 800 amino acids, depending on the length of the non-translated sequences. Likewise, the 1 kb class 20 codes for a protein of 250-300 amino acids. Enzymes that are derepressed by growth on methanol and are 700 to 800 amino acids long, most likely are MOX (Kato et al., 1976; Roa and Blobel, 1983) and DHAS (Bystrykh et al., 1981). Derepressed enzymes in the 250 to 300 amino 25 acid range are probably formaldehyde and formate dehydrogenase (Schütte et al., 1976). The polyadenylated RNA was characterized further by in vitro translation in a reticulocyte cell free translation system. Two microliters of the polyadenylated RNA directed protein 30 mixture were separated directly on a 10% SDS polyacrylamide gel, while the remaining 18 microliters were subjected to immuno-precipitation with antiserum against MOX (Fig. 7). Six strong bands dominate in the total protein mixture, having molecular weights of 35 respectively 78kd, 74kd, 58kd, 42kd, 39kd and 36kd. Essentially the same molecular weights were found by

Roa and Blobel (1983) in a total cell extract from methanol-grown H. polymorpha cells.

The 74kd protein can tentatively be assigned to the monomer of MOX, the 58kd protein to the monomer of catalase and the 39kd and 36kd proteins to the monomers of formaldehyde dehydrogenase and formate dehydrogenase, respectively. The 78kd polypeptide possibly is DHAS, while the 42kd polypeptide remains unidentified. After immuno-precipitation, both high molecular weight proteins react with the MOX antiserum.

# Cloning of the gene for MOX

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15 Although the 2.3 kb mRNA class induced by growth on methanol obviously codes for at least 2 polypeptides, it seemed a good candidate for screening a Hansenula polymorpha clone bank by hybridization. The 5-20 kb fraction of partially Sau3AI digested H. polymorpha DNA was cloned in phage lambda L47.1.

Per microgram insert DNA, 300 000 plaques were obtained while the background was less than 1:1000. Two Benton Davis blots, containing about 20 000 plaques each, were hybridized with 15 000 cpm of the mRNA-derived cDNA probe. After 3 weeks of autoradiography about 40-50 hybridizing plaques could be detected. All plaques were picked up and five were purified further by plating at lower density and by a second hybridization with the cDNA probe. From four, single hybridizing plaques (1, 3, 4, 5) DNA was isolated. The insert length varied from 8 to 13 kb.

# Hybridization selection using organic-synthetic DNA probes

The sequence of 30 amino acids at the amino terminus of

purified MOX was determined (Fig. 8).

Using the most abundant codon use for the yeast S. cerevisiae, a sequence of 14 bases could be derived 5 from part of this protein sequence, with only one ambiguity. Both probes, indicated in Fig. 4, were synthesised. In both probes an EcoRI site is present. DBM blots were made from the DNA of the MOX clones digested with the restriction enzymes BamHI, EcoRI/HindIII, 10 HindIII/SalI and PstI/SalI and separated on 1.5% agarose gels. After hybridization of the blot with a mixture of both radioactively labelled probes, the clones 1, 4 and 5 hybridize, while clone 3 does not, as shown for the HindIII/SalI blot in Fig. 9. However, the 15 probes did not hybridize with the EcoRI/HindIII digested DNA of these clones (result not shown). Since an EcoRI site is present in the probes, the hybridizing DNA in the clones probably is cut by this enzyme too. Consequently the hybridization overlap has become too 20 small to allow the formation of stable hybrids.

### Restriction map and sequence analysis

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By comparing restriction enzyme digests and by crosshybridization experiments it was concluded that clones 1, 4 and 5 covered identical stretches of DNA.

In order to definitely establish the nature of this stretch of cloned DNA the insert of clone 4 was analyzed in detail. Hybridization with the amino terminal probe showed that the complete MOX gene (ca. 2 kb) was present, including 2 kb sequences upstream and 3.5 kb downstream (Fig. 10).

DNA sequence analysis of the smallest <u>EcoRI</u> fragment revealed the nucleotide sequence corresponding to the amino terminus of MOX as was determined by amino acid sequence analysis.

For sequence analysis, several fragments were subcloned in Ml3mp8/Ml3mp9 or Ml3mpl8/Ml3mpl9 respectively in two orientations, as indicated in Fig. 10. Clones that were smaller than 0.5 kb were sequenced directly from both sides. The larger clones were cut at the unique restriction sites situated in the middle of the cloned fragment, to allow generation of exonuclease Bal31 digested subclones as described in materials and methods. Using specific oligonucleotide primers, sequences around the restriction-sites used for sub-10 cloning and sequences that did not allow an unequivocal sequence determination were sequenced once more, using the 5.5 kb BamHI/SacI subclone that covers the whole sequence. The complete nucleotide sequence is given in 15 Fig. 11A and 11B.

The sequence contains an open reading frame of 2046 nucleotides that can code for a protein of 664 amino acids. The last codon of the open reading frame codes for Phe, which is in agreement with the carboxy terminus of purified MOX. The amino acid composition derived from the DNA sequence encoding this protein, and the amino acid composition of purified MOX are virtually identical (Table III). The only important 25 differences involve the serine and threonine residues, , which are notoriously difficult to determine.

The calculated molecular weight of the protein is 74 050 Dalton, which agrees well with the molecular weight of 74 kd of MOX, as determined on polyacrylamide/SDS gels.

### Codon usage

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In Table IV the codon usage for MOX is given. A bias 35 towards the use of a selective number of codons is evident.

### Example 2

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CONSTRUCTION OF A PLASMID, PUR 3105, BY WHICH THE GENE CODING FOR NEOMYCIN PHOSPHOTRANSFERASE, THAT CONFERS RESISTANCE AGAINST THE ANTIBIOTIC G 418, IS INTEGRATED INTO THE CHROMOSOMAL MOX GENE UNDER REGIE OF THE MOX REGULON.

- H. polymorpha cells, transformed with either the plas-10 mids YEP 13, YRP 17, pHARS 1 or pHARS 2, were unstable and lost their leu+ or ura+ phenotype already after 10 generations upon growth under non-selective conditions. In order to obtain stable transformants and to test the MOX promoter, a plasmid pUR 3105 is construc-15 ted in which the neomycin phosphotransferase gene (NEO<sup>R</sup>) is brought under direct control of the MOX regulon. The construction is made in such a way that the first ATG of the NEO<sup>R</sup> gene is coupled to 1.5 kb of the MOX regulon. The cloning of such a large regu-20 lon fragment is necessary as shorter fragments, that do not contain the -1000 region of the regulon, were less efficient.
- The NEOR gene was isolated as a 1.1 kb XmaIII-SalI 25 fragment from the transposon Tn5, situated from 35 bp downstream of the first ATG up to 240 bp downstream of the TGA translational stop codon. To avoid a complex ligation mixture, first pUR 3101 is constructed (Fig. 12A), which is a fusion of the far upstream SalI-XmaIII 30 (position -1510 to position -1128) fragment of the MOX regulon, and the NEOR gene, subcloned on M13mp9. Another plasmid is constructed, pUR 3102, in which the 1.5 kb SalI-HqiAI fragment of the MOX gene, that covers nearly the whole MOX regulon, is ligated to a MOX-35 NEO<sup>R</sup> adapter (Fig. 12B) sequence and cloned in M13-mp9. The 1.2 kb XmaIII fragment of this plasmid is cloned in-

to the XmaIII site of pUR 3101, resulting in pUR 3103, which is the exact fusion of the MOX regulon and the NEOR gene (Fig. 12C). The orientation is checked by cleavage with HgiAI and SalI. From the lambda-MOX-4 clone, a SalI-SacI fragment is subcloned that reaches from the SalI site, still in the structural MOX gene (position 894), up to the SacI site, far downstream of the structural MOX gene (position 3259) (see Fig. 10). This M13mp19 subclone is called pUR 3104. The plasmid pUR 3105 is obtained by the direct ligation of the 2.7 10 kb SalI fragment from pUR 3103 into the SalI site of pUR 3104. The orientation is tested by cleavage with Smal and Sacl.

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After cleavage of this plasmid with HindIII and SacI 15 and the transformation of this cleaved plasmid to H. polymorpha, G 418-resistant colonies are found that do not lose their resistance upon growth under nonselective conditions for a large number of generations.

### Example 3

THE CONSTRUCTION OF PUR 3004, BY WHICH THE GENE CODING FOR D-AMINO ACID OXIDASE IS TRANSFERRED TO THE CHROMO-SOME OF H. POLYMORPHA UNDER REGIE OF THE MOX-REGULON

D-amino acid oxidase (AAO) is an example of an oxidoreductase for the production of which the methylotrophic H. polymorpha is extremely suited. It might be
expected that the enzyme, being an oxidase like MOX, is
translocated to the peroxisomes of the yeast that are
induced during growth on methanol or a mixture of
methanol and a fermentable sugar as carbon source and
D-amino acids as the sole nitrogen source. Under these
conditions the cell will be protected from the H<sub>2</sub>O<sub>2</sub>
produced. Alternatively, AAO can be produced without
the production of H<sub>2</sub>O<sub>2</sub>, when it is placed under
regie of the MOX- or DAS-regulon. The AAO production
will be induced by the presence of methanol in the
medium.

The amino acid sequence of the AAO enzyme has been published (Ronchi et al., 1981) and the complete gene is synthesised, using the phosphite technique (Matteuci and Caruthers, 1981). The gene is constructed in such a way that the optimal codon use for H. polymorpha, as derived from the sequence of the MOX gene, is used. Moreover, several unique restriction sites are introduced without changing the amino acid sequence, to facilitate subcloning during the synthesis. The DNA sequence is shown in Fig. 13. The gene is synthesised in oligonucleotides of about 50 nucleotides in length. Oligonucleotides are purified on 16% polyacrylamide gels. The oligonucleotides that form a subclone are added together in ligase buffer (Maniatis et al., 1982) and heated to 70°C in a waterbath. The waterbath is

slowly cooled to 16°C and  $T_4$ -ligase is added. After two hours of ligation, the DNA is separated on a 1.5% agarose gel and the fragment, having the expected length, is isolated from the gel. It is subcloned in an M13mp18 vector cleaved at the respective restriction sites situated at the end of the fragment. The gene is subcloned in this way in 4 subclones, respectively SalI-HindIII (position 39-346), HindIII-XmaI (position 346-589), XmaI-KpnI (position 589-721) and KpnI-SalI (position 721-1044). The SalI-HindIII and HindIII-XmaI subclones and the XmaI-KpnI and Kpn-I-SalI subclones are ligated together as two SalI-XmaI subclones in SalI-XmaI cleaved M13mp18. These two subclones are ligated into a SalI cleaved M13mp8, resulting in pUR 3001 (Figs 13, 14A). The whole sequence is confirmed by the determination of the nucleotide sequence using the modified Sanger dideoxy sequencing technique (Biggin et al., 1983).

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20 The construction of the integrative plasmid, containing the AAO gene is shown in Fig. 14A, B. The nearly complete AAO gene is placed upstream of the MOX termination region, by insertion of the AAO gene-containing SalI fragment of pUR 3001, in the unique SalI site of 25 pUR 3104 (see also Fig. 14A), resulting in pUR 3002. The orientation is checked by cleavage with HindIII. The MOX promoter region is isolated as a 1.4 kb SalI-HgiAI fragment from pUR 3102 (Fig. 14A). This fragment is subsequently placed upstream of the AAO gene in pUR 30 3002, by ligation to partially SalI-digested pUR 3002 in the presence of the HgiAI-SalI MOX-AAO adapter, shown in Fig. 14A. The orientation of the resulting plasmid pUR 3003 is checked again by cleavage with HindIII. This plasmid is integrated into the MOX gene 35 after cleavage with SacI and transformation to H. polymorpha cells. Transformants are selected by their ability to grow on D-amino acids as nitrogen source in

the presence of methanol as inducer.

As the selection of cells containing the AAO gene is not simple, another selection marker is introduced. To this end, the S. cerevisiae LEU2 gene is integrated 5 in between the structural AAO gene and the MOX terminater. For this construction, the plasmid pURS 528-03 is used. This plasmid is derived from pURY 528-03 described in European patent application 0096910. The construction is shown in Fig. 14C. The 10 deleted carboxy terminal LEU2 gene sequence of pURY 528-03 was replaced by the complete carboxy terminal LEU2 gene sequence from pYeleu 10 (Ratzkin and Carbon, 1977) and the E. coli lac-lac regulon was eliminated. Subsequently the HpaI-SalI fragment of pURS 528-03 15 containing the LEU2 gene, is blunt end inserted in the SalI site of pUR 3003, situated in between the AAO structural gene and the MOX terminater. The orientation of the resulting plasmid pUR 3004 can be checked by cleavage with SalI and SacI. pUR 3004 integrates in the 20 chromosomal MOX gene of H. polymorpha after transformation of the SacI-cleaved plasmid to a H. polymorpha leu mutant. Selected leu transformants are integrated in the chromosomal MOX gene, together with the AAO gene. 25

# Example 4

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THE CONSTRUCTION OF PUR 3204, PUR 3205, PUR 3210 and PUR 3211, BY WHICH THE SMALL PEPTIDE HORMONE, THE HUMAN GROWTH RELEASING FACTOR, IS EXPRESSED UNDER REGIE OF THE MOX-REGULON, EITHER BY INTEGRATION INTO THE CHROMOSOMAL MOX GENE (PUR 3203, PUR 3204), OR BY INTEGRATION INTO A HARS1-CONTAINING PLASMID (PUR 3205) OR BY FUSION TO THE MOX STRUCTURAL GENE (PUR 3209, PUR 3210 and PUR 3211).

Human growth hormone releasing factor (HGRF) is a small, 44 amino acids long, peptide, that activates the secretion of human growth hormone from the pituitary 15 glands. HGRF can be used in the diagnosis and treatment of pituitary dwarfism in man. Since HGRF has been shown to induce growth hormone stimulation in numerous species, HGRF might be used in the vetinary field too, by stimulating growth of animals and increase of milk 20 production (Coude et al., 1984). It is difficult to obtain HGRF from human sources, but it could very well be produced by biotechnological processes, once the gene has been cloned and transferred to an appropriate host organism. Also, as a general example of the production 25 of a peptide hormone by  $\underline{H}$ .  $\underline{polymorpha}$ , the gene for HGRF is synthesised in the optimal codon use of H. polymorpha and brought to expression in several ways.

For the construction of pUR 3204 and pUR 3205, the gene fragment that codes for the carboxy terminal part of the protein is synthesised in DNA oligomers of about 50 nucleotides in length and subcloned as a <a href="https://mindill-SalI"><u>HindIII-SalI</u></a> fragment in <a href="https://mindill-SalI"><u>HindIII-SalI</u></a> cleaved M13mp18, resulting in pUR 3201 (Figs 15, 16A). This <a href="https://mindill-SalI"><u>HindIII-SalI</u></a> fragment is subsequently inserted upstream of the MOX terminater in <a href="https://mindill-SalI"><u>HindIII-SalI</u></a> cleaved pUR 3104 (Fig. 16A), resulting in

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pUR 3202. The MOX promoter is inserted in front of the HGRF gene, by insertion of the SalI-HgiAI MOX-promoter fragment from pUR 3102 (Fig. 16A) in HindIII cleaved pUR 3202, using a HgiAI-HindIII adapter between the MOX-promoter and the HGRF gene (Figs 15, 16A). The orientation of the resulting plasmid pUR 3203 is checked by cleavage with SalI and HgiAI. pUR 3203 integrates into the chromosomal MOX gene of H. polymorpha after transformation of the SacI cleaved plasmid. Transformants are selected on immunological activity. pUR 3203 is cleaved with SalI, to insert the SalI-HpaI fragment of pURS 528-03 (Fig. 16B) that contains the LEU2 gene. The orientation of this gene in pUR 3204 is checked by cleavage with HindIII and EcoRI. pUR 3204 integrates into the chromosomal MOX gene of H. polymorpha after transformation of the SacI cleaved plasmid (Fig. 16B) to a leu H. polymorpha mutant. Selection on on leu<sup>+</sup> transformants. A plasmid, called pUR 3205, that replicates autonomously in H. polymorpha and contains the HGRF gene, is obtained by insertion of the EcoRI, partially HindIII cleaved 4 kb long fragment of pUR 3203, containing the HGRF gene inserted in between the MOX-promoter and terminater, into partially HindIII-EcoRI cleaved pHARS1 (Figs 2, 16C). The construction of pUR 3205 is checked by cleavage with HindIII.

The production of small peptides as HGRF by microorganisms is often unstable as a result of enzymic
degradation (Itakura et al., 1977). Fusion to a protein
like MOX, and subsequent transport to the peroxisomes,
could prevent degradation. Therefore, we decided to insert the HGRF gene into the unique KpnI site at
position 1775 (amino acid 591, Figs 10, 11) of the MOX
structural gene. The HGRF gene is synthesised again in
DNA oligomers of 50 nucleotides in length, but now as
two KpnI-HindIII subclones that are cloned as a complete HGRF structural gene in M13mp19, cleaved with

KpnI (plasmid pUR 3206, Figs 17, 16D). Moreover, the ATG triplet coding for the internal methionine of HGRF at position 27 (Coudé et al., 1984) (position 82 of the DNA sequence) is converted into a TGT triplet coding 5 for cysteine. This does not alter the HGRF activity essentially, and facilitates the cleavage of HGRF from the fusion protein by CNBr cleavage (Itakura et al., 1977). From phage lambda MOX-4 (Fig. 10 SphI (position -491)-KpnI fragment is isolated and in 10 serted into SphI-KpnI cleaved Ml3mpl9. This results in pUR 3207. pUR 3206 is cleaved with KpnI and the HGRF gene is inserted into the KpnI site of pUR 3207, resulting in pUR 3208. The orientation is checked by direct sequence analysis on the single-stranded DNA of 15 pUR 3208. Subsequently the downstream part of the MOX gene, from the unique KpnI site up to the SacI site, is isolated as a 1.5 kb fragment from phage lambda MOX-4 and inserted into SacI - partially KpnI cleaved pUR 3208. The orientation of the resulting plasmid pUR 20 3209 is checked by digestion with KpnI. pUR 3209 integrates into the chromosomal MOX gene of H. polymorpha after transformation of the SacI, SphI cleaved plasmid. Selection on immunological activity.

25 This MOX-HGRF fusion gene is inserted into pHARS1 by isolation of the whole fusion gene from partially HindIII, partially EcoRI cleaved pUR 3209, into EcoRI partially HindIII cleaved pHARS1. This results in pUR 3210, which replicates in H. polymorpha after trans-30 formation (Fig. 16E). Alternatively, the LEU2containing SalI-HpaI fragment of pURS 528-03 is inserted into the blunt-ended KpnI site of the HGRF gene, located at the carboxy terminus of the encoded protein, after partial KpnI cleavage of pUR 3209. The resulting 35 plasmid pUR 3211 integrates into the chromosomal MOX gene of H. polymorpha, after transformation of the SacI, SphI cleaved plasmid (Fig. 16F).

### Discussion

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From the length of the open reading frame, from the similarity in the amino acid composition of purified MOX and the DNA derived protein sequence and from the identical 30 N-terminal amino acids, it is concluded that the complete gene for MOX from the yeast Hansenula polymorpha has been cloned. Its calculated molecular weight agrees well with the molecular weight determined on SDS polyacrylamide gels. Apart from the coding sequence, more than 1200 bp has been sequenced from both the 5'- and the 3'-non-coding regions, reaching from the SalI site upstream of the coding sequence, up to the SacI site downstream. The gene appears not to be interrupted with intervening sequences.

The protein is not transcribed in the form of a precursor. Based on the determination of the molecular weight, N-terminal signal sequences could not be detected in earlier studies of Roa and Blobel (1983) or Roggenkamp et al. (1984) as well. In similar studies, it was suggested that also the rat liver peroxisomal enzymes uricase (Goldman and Blobel, 1978) and catalase (Goldman and Blobel, 1978; Robbi and Lazarow, 1978) do not contain a cleavable N-terminal signal peptide. However, as discussed by these authors, proteolytic degradation could possibly explain the lack of the detection of such a signal sequence.

Our sequence results definitely prove that for translocation of this protein to the peroxisome, a cleavable
N-terminal signal sequence is not required. Such a
translocation signal may well be situated in the
internal sequence of the mature protein, as is the case
for ovalbumine (Lingappa et al., 1979). Inspection of
the protein sequence reveals the amino acid sequence
Gly X Gly Y Z Gly (amino acids 13-18), which is charac-

teristic for FAD-(flavin adenine dinucleotide)-containing enzymes (Ronchi et al., 1981).

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The isolation of the MOX gene described above gives a way how to determine the DNA sequence coding for MOX and the amino acid sequence of the MOX enzyme.

Similarly, the DNA sequences and amino acid sequences belonging to other oxidase-enzymes can be isolated and determined. The knowledge of the MOX gene sequence can be used to facilitate the isolation of genes coding for alcohol oxidases or even other oxidases. By comparing the properties and the structure of enzymes one can probably establish structure function and activity relationships. One can also apply methods as site-directed mutagenesis, or shortening or lengthening of the protein coding sequences, modifying the corresponding polypeptides, to select oxidase-enzymes with improved properties, e.g. with increased alkali stability, improved production, or oxidase-enzymes which need a substrate which is more compatible with detergent products.

Besides the isolation and characterization of the

structural gene for MOX from the yeast H. polymorpha,
also the isolation and characterization of the structural gene for DHAS from the yeast H. polymorpha has
been carried out in a similar way.

The DNA sequence of DAS is given in Fig. 18A-18C. A restriction map is given in Fig. 19. The amino acid composition calculated from the DNA sequence of DAS appeared to be in agreement with the amino acid composition determined after hydrolysis of purified DHAS.

The DHAS enzyme catalyses the synthesis of dihydroxyacetone from formaldehyde and xylulose monophosphate. This reaction plays a crucial role in the methanol-

assimilation process (cf. Veenhuis et al., 1983).

As described before, the synthesis of MOX and DHAS is subject to glucose repression. It has now been found that higher levels of MOX are reached when using glucose/methanol mixtures as substrates instead of 0.5% (v/v) methanol. Under the former conditions up to 30% of the cellular protein consists of MOX, compared with up to 20% under the latter conditions.

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It was considered that in the regulons of MOX and DAS sequences must exist that play a decisive role in the regulation of repression/derepression by glucose or of the induction by methanol. Some homology therefore might be expected.

A striking homology of the "TATA-boxes" has been found, both having the sequence CTATAAATA. No other homologies in the near upstream region of the MOX and DAS regulons have been found. Unexpectedly, a detailed study of both regulons has shown a remarkable homology of the regulons for MOX and DAS in the region about 1000 bp upstream of the translation initiation codon. A practically complete consecutive region of 65 bp in the regulon of MOX is homologous to a 139 bp region in the DAS regulon, interspersed by several non-homologous regions (see Fig. 20). A similar homology is not found in any other region of both genes, that are over 4 kb in length including their upstream and downstream sequences. It is suggested that these homologous sequences play a role in the regulation of both genes by glucose and methanol. Transformation studies with vectors containing as regulon the first 500 bp upstream of the ATG of the structural gene of MOX, showed that this shortened MOX-regulon gave rise to a relatively low expression of the indicator gene beta-lactamase. Indicator genes are genes which provide the yeast with

properties that can be scored easily, e.g. the gene for neomycin phosphotransferase giving resistance to the antibiotic G 418 (cf. Watson et al., 1983) or an auxotrophic marker such as leucin.

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The fact that the far upstream homologous regions in the MOX and DAS genes have different interruptions and the fact that DAS is repressed at 0.1% glucose and MOX is not, suggest that these homologous regions are of importance to the repression-derepression by glucose and/or the induction of the expression in the presence of methanol. This assumption has been found correct indeed, and the presence or absence of these homologous regions can therefore be important for specific applications. For example, if the -1052 to -987 region of the MOX gene or the -1076 to -937 region of the DAS gene is important for the induction of MOX or DAS by methanol, the presence of these regions is required for the expression of MOX or DAS and/or for the induction of other enzymes by methanol. Another example might be the removal of the regions to avoid repression by glucose, which is needed for the expression of genes coding for proteins other than MOX and DHAS under influence of the MOX and/or DAS regulatory regions with glucose as a carbon source.

Thus one aspect of the present invention relates to the isolation and complete characterization of the structural genes coding for MOX and DHAS from the yeast H. polymorpha. It further relates to the isolation and complete characterization of the DNA sequences that regulate the biosynthesis of MOX and DHAS in H. polymorpha, notably the regulons and terminaters.

Moreover, it relates to combinations of genes coding for alcohol oxidase or other oxidases originating from H. polymorpha strains other than H. polymorpha CBS

4732, or <u>Hansenula</u> species other than <u>H. polymorpha</u>, or yeast genera other than <u>Hansenula</u>, or moulds, or higher eukaryotes, with the powerful regulon and terminater of the MOX gene from <u>H. polymorpha</u> CBS 4732. These combinations may be located on vectors carrying amongst others an autonomously replicating sequence originating from <u>H. polymorpha</u> or related species or minichromosomes containing centromers, and optionally selection marker(s) and telomers. These combinations may also be integrated in the chromosomal DNA of <u>H. polymorpha</u>.

Furthermore it relates to combinations of the powerful regulon or parts of it and terminaters of the MOX and/or DAS and - by site-directed mutagenesis or other methods - changed structural genes coding for alcohol oxidase or another oxidase. These changed structural genes may be located on episomal vectors, in minichromosomes or integrated in the chromosomes of H. polymorpha, H. wingeii, H. anomala, and S. cerevisiae or in other yeasts.

Besides this, the present invention relates to combinations of the regulon and terminater of the MOX and/or DAS gene of  $\underline{H}$ .  $\underline{polymorpha}$  with structural genes coding for other proteins than oxidases.

A very important and preferred embodiment of the invention is a process for preparing a polypeptide, such as a protein or an enzyme, by culturing a microorganism under suitable conditions, optionally concentrating the polypeptide and collecting same in a manner known perse, characterized in that a microorganism is used that has been obtained by recombinant DNA technology and caries a structural gene coding for the polypeptide concerned, the expression of which is under the control of a regulon, comprising a promoter and at least either the -1052 to -987 region of the MOX gene of Hansenula

polymorpha CBS 4732, or the -1076 to -937 region of the DAS gene of <u>Hansenula polymorpha</u> CBS 4732, or a corresponding region of other methylotrophic moulds or yeasts, or an effective modification of any of these regions.

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Surprisingly, it has been observed by the present inventors that the regions concerned, which are shown in Fig. 20 and are referred to herein as the -1000 regions of the MOX and DAS genes, are of crucial importance for the expression of the structural gene concerned. Experiments performed with recombinants containing the MOX regulon from which this region was eliminated showed a low level of expression. Therefore, use of a regulon comprising such -1000 region, or an effective modification thereof, i.e. any modification which does not result in a significant mutilation of the function of said region, makes it possible to realize production of a relatively high amount of the desired polypeptide.

A preferred embodiment of this process according to the invention is characterized in that the structural gene concerned has been provided with one or more DNA sequences coding for amino acid sequences involved in the translocation of the gene product into the peroxisomes or equivalent microbodies of the microbial host. Translocation of the produced polypeptide into the peroxisomes or equivalent microbodies improves their stability, which results in a higher yield. For certain kinds of polypeptides, in particular oxidases, such translocation is imperative for survival of the microbial host, i.e. to protect the host against the toxic effects of the hydrogen peroxide produced when the microbial host cells are growing on the substrate of the oxidase. If the oxidase concerned does not contain addressing signals which are functional in the microbial host used in the production process, one

should provide the structural gene with sequences coding for host specific addressing signals, for example by adding such sequences or by substituting these for the original addressing sequences of the gene. Production of a fused polypeptide, in which the fusion partner carries suitable addressing signals, is another possibility. In case methylotrophic yeasts are used in the production process, it is preferred that the DNA sequences consist of the MOX gene or thos parts thereof which are responsible for MOX translocation into the peroxisomes or microbodies.

Finally, this aspect of the present invention is related to the synthesis of MOX originating from  $\underline{H}$ .  $\underline{poly}$ -morpha in other yeasts.

Some microorganisms with the potential of producing alcohol oxidases are summarized below.

Yeasts producing alcohol oxidases
(Taxonomic division according to Lee and Komagata, 1980)

Group 1 Candida boidinii

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Group 2a Hansenula philodendra

Pichia lindnerii

Torulopsis nemodendra

" pinus

sonorensis

	Group 2b	Candida cariosilignicola					
		Hansenula glucozyma					
		" henricii					
		" minuta					
5		" nonfermentans					
		" polymorpha					
		" wickerhamii					
		Pichia pinus					
		" trehalophila					
10							
	Group 2c	Candida succiphila					
		Torulopsis nitratophila					
	Group 3	Pichia cellobiosa					
15	Q.,	Wanaanula aanaulaka					
	Group 4	Hansenula capsulata					
		Pichia pastoris Torulopsis molischiana					
		TOTATOPSIS MOTISCHIANA					
20	Moulds pr	oducing alcohol oxidases:					
		Lenzites trabea					
		Polyporus versicolor					
		" obtusus					
		Poria contigua					
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	Among the oxidases other than alcohol oxidases, the						
	most interesting are:						
	- glycerol oxidase,						
	- ald	ehyde oxidase,					
30		ne oxidase,					
	<ul><li>aryl-alcohol oxidase,</li></ul>						
		no acid oxidase,					
	_	cose oxidase,					
2.5	_	actose oxidase,					
35		bose oxidase,					
		c acid oxidase, oroperoxidase, and					

- xanthine oxidase.

Combinations of the powerful regulons and terminaters of the MOX and DAS genes from H. polymorpha and structural genes for oxidases may be combined with one or more DNA sequences that enable replication of the structural gene in a particular host organism or group of host organisms, for example autonomously replicating sequences or centromers (and telomers) originating from H. polymorpha, to suitable vectors that may be transferred into H. polymorpha and related yeasts or other microorganisms.

H. polymorpha mutants LEU-1 and LR9, mentioned on page 12 of this specification, were deposited at the Centraalbureau voor Schimmelcultures at Delft on 15th July, 1985, under numbers CBS 7171 and CBS 7172, respectively.

The above description is followed by a list of references, claims, Tables, Legends to Figures and Figures.

#### References

- 1. GB-PS 1 225 713 (Colgate-Palmolive Company; publ. 24th March 1971; priority date 19th April 1968).
- 2. DE-PA 2 557 623 (Henkel & Cie GmbH; publ. 30th June 1977; priority date 20th December 1975).
- 3. GB-PA 2 101 167 (Unilever PLC; publ. 12th January 1983; priority date 7th July 1981).
  - van Dijken, J.P., Otto, R. and Harder, W. (1976),
     Arch.Microb. <u>111</u>, 137-144.
- 5. Veenhuis, M., van Dijken, J.P. and Harder, W. (1983) in Advances in Microbial Physiology, Rose, A.H., Gareth Morris, J. and Tempest, D.W., Eds, Vol. 24, pp 1-82, Academic Press, New York.
- 6. Roggenkamp, R., Janowicz, Z., Stanikowski, B. and Hollenberg, C.P. (1984), Mol.Gen.Genet. 194, 489-493.
- 7. Sahm, H. (1977) in Advances in Microbiol. Engineering, Ghose, T.K., Fiechter, A. and Blakebrough,
  N., Eds, Vol. 6, pp 77-103, Springer-Verlag,
  Berlin.
- 8. Bystrykh, L.V., Sokolov, A.P. and Trotsenko, Y.A. (1981), FEBS Letters <u>132</u>, 324-328.
  - Roa, M. and Blobel, G. (1983), Proc. Natl. Acad. Sci. USA, 80, 6872-6876.
- 35 10. Veenhuis, M., van Dijken, J.P., Pilon, S.A.F. and Harder, W. (1978), Arch. Microbiol. <u>117</u>, 1953-163.

- 11. Loenen, W.A.M. and Brammar, W.J. (1980), Gene 20, 249-259.
- 12. Hohn, B. (1979) in Methods in Enzymology, Wu, R., Ed., Vol. 68, pp 299-309, Academic Press, New York.
- 13. Edens, L., Heslinga, L., Klok, R., Ledeboer, A.M.,
  Maat, J., Toonen, M.Y., Visser, C. and Verrips, C.T.

  (1982), Gene 18, 1-12.
  - 14. Pelham, H.R.B. and Jackson, R.J. (1976), Eur.J. Biochem. 67, 247-257.
- 15 15. Valerio, D., Duyvensteijn, M.G.C., Meera Khan, P., Geurts van Kessel, A., de Waard, A. and van der Eb, A.J. (1983), Gene 25, 231-240.
- 16. Ledeboer, A.M., Verrips, C.T. and Dekker, B.M.M. (1984), Gene 30, 23-32.

- 17. Maniatis, T., Fritsch, E.F. and Sambrook, J. (1982), Molecular Cloning, p 278, Cold Spring Harbor Laboratory Publ., New York.
- 18. Benton, W.D. and Davis, R.W. (1977), Science 196, 180-182.
- 19. Ambler, R.P. (1972), Methods in Enzymology, Vol. 25, pp 262-272, Academic Press, New York.
  - 20. Matteucci, M.D. and Caruthers, M.H. (1981), J.Am. Chem. Soc. 103, 3185-3191.
- 35 21. Wallace, R.B., Johnson, M.J., Hirose, T., Miyake, T., Kawashima, E.H. and Itakura, K. (1981), Nucl. Acids Res. 9, 879-894.

- 22. Biggin, M.D., Gibson, T.J. and Hong, G.F. (1983), Proc.Natl.Acad.Sci.USA 80, 3963-3965.
- Gleeson, M.A., Waites, M.J. and Sudbery, P.E. (1984),
   in: Microbial growth on C<sub>1</sub> compounds, Eds.
   Crawford, R.L. and Hanson, R.S., Publ. A.S.M.,
   Washington, 228-235.
- 24. Fink, G.D. (1970), Methods in Enzymology, Tabor, H.
  and Tabor, C.W., Eds., Vol. 17, pp 59-78, Academic Press, New York.
  - 25. Boeke, J.D., LaCroute, F. and Fink, G.D. (1984), Mol. Gen.Genet. <u>197</u>, 345-346.
  - 26. Jones, E.W. and Fink, G.D. (1982), Cold Spring Harbour Monogr.Ser., 11B, 181-299.

- 27. Lieberman, I., Kornberg, A. and Simms, E.S. (1955),
   J.Biol.Chem. 215, 403-415.
  - 28. Stinchcomb, D.T., Thomas, M., Kelly, J., Selker, E. and Davis, R.W. (1980), Proc.Natl.Acad.Sci.USA 77, 4559-4563.
- 29. Stinchcomb, D.T., Mann, C. and Davis, R.W. (1982), J.Mol.Biol. <u>158</u>. 157-179.
- 30. Struhl, K., Stinchcomb, D.T., Scherer, S. and Davis, R.W. (1979), Proc.Natl.Acad.Sci.USA 76. 1035-1039.
  - 31. Carlson, M. and Botstein, D. (1982), Cell 28, 145-154.
- 35 32. Beggs, J.D. (1978), Nature 275, 104-109.
  - 33. Broach, J.R., Strathern, J.N. and Hicks, J.B. (1979), Gene 8, 121-133.

- 34. Das, S., Kellerman, E. and Hollenberg, C.P. (1984), J.Bacteriol. 158, 1165-1167.
- 35. Ito, H., Fukuda, Y., Murata, K. and Kimura, A. (1983), J.Bacteriol. <u>153</u>, 163-168.
  - 36. Klebe, R.J., Harriss, J.V., Sharp, Z.D. and Douglas, M.G. (1983), Gene 25, 333-341.
- 37. Hollenberg, C.P. (1982), Curr.Top.Microbiol.Immunol. 96, 119-144.
  - 38. Kato, N., Omori, Y., Tani, Y. and Ogata, K. (1976), Eur.J.Biochem. 64, 341-350.
- 39. Schütte, H., Flossdorf, J., Sahm, H. and Kula, M.R., (1976), Eur.J.Biochem. 62, 151-160.

- 40. Ronchi, S., Minchiotti, L., Galliano, M., Curti, B.,
  Swenson, R.P., Williams, C.H. and Massey, V. (1981),
  J.Biol.Chem. 257, 8824-8830.
  - 41. Ratzkin, B. and Carbon, J. (1977), Proc.Natl.Acad. Sci.USA 74, 487-491.
  - 42. Coudé, F.X., Diaz, J., Morre, M., Roskam, W. and Roncucci, R. (1984), Trends in Biotechnology 2, 83-88.
- 30 43. Itakura, K., Hirose, T., Crea, R., Riggs, A.D., Heyneker, H.L., Bolivar, F. and Boyer, H.W. (1977), Science 198. 1056-1063.
- 44. Goldman, B.M. and Blobel, G. (1978), Proc.Natl. Acad.Sci.USA 75, 5066-5070.
  - 45. Robbi, M. and Lazarow, P.B. (1978), Proc.Natl.Acad. Sci.USA 75, 4344-4348.

- 46. Lingappa, V.R., Lingappa, J.R. and Blobel, G. (1979), Nature 281, 117-121.
- 47. Watson, J.D., Tooze, J. and Kurtz, D.T. (1983),

  Recombinant DNA, A Short Course, page 178, published
  by W.H. Freeman and Company, New York.
  - 48. Lee, J.D. and Komagata, K. (1980), J.Gen.Appl. Microbiol. <u>26</u>, 133-158.

TABLE I

Activities of orotidine 5'-phosphate decarboxylase and orotidine 5'-phosphate pyrophosphorylase in <u>H. poly-</u>morpha mutants requiring uracil for growth.

Strain/	Reversion	Activity (%) <sup>a</sup>		
Genotype	rate	Orotidine 5'- phosphate decarboxylase	Orotidine 5- phosphate pyrophosphorylase	
Wild type	_	100	100	
LR 9/odcl	<2 × 10 <sup>9</sup>	<b>&lt;</b> 1	106	
MR 7/odcl	6 × 10 <sup>7</sup>	<b>\ 1</b>	71	
NM 8/odcl	$3 \times 10^{8}$	<b>لا</b>	105	
CLK 55/opp	l n.e.b	90	<1	
CLK 68/opp		82	<b>&lt;</b> 1	
YNN 27/ura		0	n.e.	

Strains were grown in YEPD until late exponential phase. Extraction of cells was performed with glass beads using a Braun homogenizer. Protein was estimated by the optical density at 280 nm.

a) Expressed as the percentage of wild type activity.

<sup>30</sup> b) Not estimated.

TABLE II

Transformation of uracil-requiring mutants of  $\underline{H}$ .  $\underline{poly}$ -morpha

Strain	Plasmid	Transformation frequency <sup>a</sup>	Stability <sup>b</sup> (%)	Status ( transform DNA
LR 9	YRP17	2.2 x 10 <sup>2</sup>	<b>〈</b> 1	Autonomo replicat
LR 9	pHARS1	$1.5 \times 10^3$	2	Autonomo replicat
LR 9	pHARS2	$4.6 \times 10^2$	1.5	Autonomo replicat
LR 9	YIP5	3 (38) <sup>C</sup>	105	Integrat
LR 9	pRB58	0	_	-
LR 9	рнн85	0	-	-
YNN 27	YIP5	0	_	_

a) Expressed as total number per /ug of DNA. Intact cells treated with polyethyleneglycol were used for transformation as described in Materials and Methods.

b) Expressed as the percentage of remaining uracil prototrophs after growth on YEPD for ten generations.

Number in parentheses indicates the amount of minicolonies containing free plasmid YIP5.

TABLE III

### Amino acid composition of MOX

5	Amino	Acid	DNA	sequence	Hydro	lysate	a)
	Dun	•		31		32	
	PHE						
	LEU			47		49	
	ILE			34		34	
10	MET			12		11	
	VAL			42		43	
	SER			43		33 a)	
	PRO			43		42	
	THR			44		38	
15	ALA			47		50	
	TYR			27		27	
	HIS			19		21	
	GLN			13	_		
	GLU			36	]	51	
20	ASN			32			
	· ASP			50	.J	84	
	LYS	•		35		<b>3</b> 8	
	CYS			13		12	
	TRP			10		_ b)	
25	ARG			36		36	
	GLY			50		53	

a) Hydrolysis was performed for 24 h.

<sup>30</sup> b) Not determined.

TABLE IV

Comparison of preferred codon usage in <u>S</u>. <u>cerevisiae</u>,

H. polymorpha and <u>E</u>. <u>coli</u>

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		Saccha	romyces	Hansenula	E. coli
				MOX	
	ALA	GCU,	GCC	GCC	GCC not used,
					no clear pref.
10	SER	UCU,	UCC	UCC, UCG	ucu, ucc
	THR	ACU,	ACC	ACC	ACU, ACC
	VAL	GUU,	GUC	GUA not used,	GUU, GUA
				no clear pref.	
	ILE	AUU,	AUC	AUC, AUU	AUC
15	ASP	GAC		GAC	GAC
	PHE	UUC		UUC	UUC
	TYR	UAC		UAC	UAC
	CYS	UGU		no clear pref.	no clear pref.
	ASN	AAC		AAC	AAC
20	HIS	CAC		CAC	CAC
	GLU	GAA		GAG	GAA
	GLY	GGU		GGC practically	
			not	t used, no clear pro	ef.
	GLN	CAA		CAG	CAG
25	LYS	AAG		AAG	AAA
	PRO	CCA		CCU, CCA	CCG
	LEU	UUG		CUG, CUC	CUG
	ARG	AGA		AGA	CGU

#### Legends to Figures

- Fig. 1. The exonuclease Bal31 digestion strategy used in sequencing specific MOX subclones. The fragment X-Y subcloned in M13mp-8 or -9, -18 or -19 5 is cut at the unique restriction site Z. The DNA molecule is subjected to a time-dependent exonuclease Bal31 digestion. The DNA fragment situated near the M13 sequencing primer is removed using restriction enzyme Y; ends are 10 made blunt end by incubation with  $T_A$ -DNA polymerase and then ligated intramolecularly. Phage plaques are picked up after transformation and the fragment is sequenced from site Z in the direction of site X. Using the 15 M13 derivative with a reversed multiple cloning site, the fragment is sequenced from site Z in the direction of site X.
- 20 Fig. 2. Alignment of pHARS plasmids derived by insertion of HARS fragments into the single SalI site of YIp5.
- Fig. 3. The complete nucleotide sequence of the HARS-1 fragment.
- Fig. 4. Estimation of copy number by Southern hybridization of H. polymorpha transformants. An
  aliquot of 8 and 16 /ul of each probe was
  electrophoresed. Lane 1, phage lambda DNA digested with HindIII and EcoRI. Lanes 2,3 transformant of K. lactis containing two copies of
  integrated plasmid, digested with HindIII (M.
  Reynen, K. Breunig and C.P. Hollenberg, unpublished); lanes 4-7, YNN 27, transformed with
  pRB58 (4-5) and YRP17 (6-7) digested with EcoRI
  respectively; lanes 8,9, LR9 transformed with

YRP17 digested with EcoRI; lanes 10,11, LR9 transformed with pHARS2 digested with HindIII; lanes 12,13, LR9 transformed with pHARS1 digested with EcoRI.

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Fig. 5. Autoradiogram of Southern blots of DNA from H. polymorpha mutant LR9 transformed by integration of plasmid YIp5. Lane 1, phage lambda DNA, digested both with HindIII and EcoRI; lane 2, pHARS-1, undigested; lanes 3-5 and lanes 6,7 show DNA from 2 different transformants. Lane 3, undigested; lane 4, digested with EcoRI; lane 5, digested with PvuII; lane 6, digested with EcoRI; lane 7, digested with PvuII; lane 8, plasmid YIp5, digested with EcoRI. Nicktranslated YIp5 was used as a hybridization probe.

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Fig. 6 Electrophoresis of <sup>32</sup>p-labelled RNA from Hansenula polymorpha, purified once (lane A) or twice (lane B) on oligo(dT)cellulose. Electrophoresis was performed on a denaturing 7 M urea 2.5% polyacrylamide gel. The position of the yeast rRNA's and their respective molecular 25 weights are indicated by 18S and 25S. The 2.3 kb band, that can be seen in lane B, was converted into a cDNA probe which was subsequently used to isolate MOX and DHAS clones from the Hansenula polymorpha clone bank.

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35S-labelled proteins obtained after in vitro Fig. 7 translation of methanol derepressed, Hansenula polymorpha mRNA with a rabbit reticulocyte lysate. Either 2 microliters of the total lysate (lane A) or an immuno-precipitate of the 35 remaining 18 microliters using a MOX specific antiserum (lane B) were separated on an 11.5%

SDS-polyacrylamide gel. A mixture of proteins with known molecular weights was used as markers.

Fig. 8. The N-terminal sequence of purified MOX, as determined on a Beckman sequenator. The two probes that could be derived from the sequence Pro-Asp-Gln-Phe-Asp, using Saccharomyces preferred codons, are indicated.

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- Fig. 9. Hybridization of a DBM blot of HindIII/SalI cut MOX clones. The DNA was separated on a 1.5% agarose gel (Fig. 9A) and the blot was hybridized to a mixture of both MOX-derived synthetic DNA probes (Fig. 8). Only one band of clones 1, 4 and 5 hybridize (Fig. 9B), indicated by an arrow in Fig. 9A. Lane M: molecular weight markers as indicated. Lane A, B, C and D: clones 1, 3, 4 and 5, respectively. Lane E: lambda L47.1.
- Fig. 10. Restriction map for MOX clone 4. Only relevant restriction sites are indicated that have been used for subcloning and sequencing of the MOX gene. The open reading frame, containing the structural MOX sequence, and the M13 subclones made are depicted. Restriction sites used are:

  B= BamHI, E<sub>I</sub> = EcoRI, E<sub>V</sub> = EcoRV, P = PstI, S1 = SalI, Sc = SacI, St = StuI, H = HindIII, Sp = SphI, K = KpnI, Hg = HgiAI and X = XmaI.
  - Fig. 11A,B. The nucleotide sequence of the MOX structural gene and its 5'- and 3'-flanking sequence.
  - Fig. 12A,C. The construction of plasmid pUR 3105 by which the neomycin phosphotransferase gene

integrates into the chromosomal MOX gene of H. polymorpha.

- Fig. 12B. Promoter MOX-neomycin phosphotransferase adapter fragments.
- Fig. 13. The DNA sequence of the AAO gene, derived from the published amino acid sequence. The gene is synthesised in the optimal codon use for

  H. polymorpha in oligonucleotides of about 50 nucleotides long. Restriction sites, used for subcloning are indicated. The HgiAI-SalI fragment forms the adapter between the structural AAO gene and the MOX promoter. The translational start codon (met) and stop codon (\*\*\*) are indicated. The structural sequence is numbered from 1 to 1044, while the MOX promoter is numbered from -34 to -1.
- 20 Fig. 14A. The construction of pUR 3003, by which the AAO gene integrates into the chromosomal MOX gene of <u>H. polymorpha</u>. Selection on activity of the AAO gene.
- 25 Fig. 14B. The construction of pUR 3004, by which the AAO gene integrates into the chromosomal MOX gene of a <u>H. polymorpha</u> leu derivative.

  Selection on leu+.
- 30 Fig. 14C. The construction of pURS 528-03. Owing to the removal of the pCR1 sequence and the double lac UV5 promoter, this plasmid is about 2.2 kb shorter than pURY 528-03.
- 35 Fig. 15. The DNA sequence of the HGRF gene, derived from the published amino acid sequence. The gene is synthesised in the optimal codon use

for <u>H. polymorpha</u> in oligonucleotides of about 50 nucleotides long. <u>HgiAI</u>, <u>HindIII</u> and <u>SalI</u> sites are used for subcloning. The <u>HgiAI</u>
<u>HindIII</u> fragment forms the adapter between the structural HGRF gene and the MOX promoter. The translational start codon (met) and stop codon (\*\*\*) are indicated. The structural sequence is numbered from 1 to 140, while the MOX promoter is numbered from -34 to -1.

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Fig. 16A. The construction of pUR 3203, by which the gene coding for HGRF integrates into the chromosomal MOX gene of <u>H. polymorpha</u>.

Selection on immunological activity of HGRF.

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Fig. 16B. The construction of pUR 3204, by which the gene coding for HGRF integrates into the chromosomal MOX gene of a <u>H. polymorpha</u> leuderivative. Selection on leude.

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Fig. 16C. The construction of pUR 3205, by which the gene coding for HGRF is inserted into a HARS-1-containing plasmid, which replicates autonomously in <u>H. polymorpha</u>. Selection by transformation of a ura mutant.

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Fig. 16D. The construction of pUR 3209, by which the gene coding for HGRF integrates into the chromosomal MOX gene of H. polymorpha, fused to the structural MOX gene. HGRF is cleaved from the fusion protein by CNBr cleavage. Selection on immunological activity of HGRF.

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Fig. 16E. The construction of pUR 3210, by which the gene coding for HGRF is inserted into a HARS-1-containing plasmid, fused to the structural MOX gene. Selection as in Fig. 16C.

Fig. 16F. The construction of pUR 3211, by which the gene coding for HGRF integrates into the chromosomal MOX gene of a <u>H. polymorpha leuderivative</u>, fused to the structural MOX gene. Selection on leut.

- Fig. 17. The DNA sequence of the HGRF gene, derived from the published amino acid sequence. The gene is synthesised as mentioned in Fig. 15, but constructed in such a way that it could be inserted into the unique KpnI site of the structural MOX gene. Therefore it was equipped with KpnI sites on both sides of the gene, and KpnI-HindIII fragments were used for subcloning. Synthesis will be as a fusion product to the MOX enzyme. The internal met (ATG) at position 82 is converted into a cys (TGT). Translational start (met) and stop (\*\*\*) codons are indicated.
- Fig. 18A,B,C. The nucleotide sequence of the DAS structural gene and its 5'- and 3'-flanking sequence.
- 25 Fig. 19. Restriction map for the DAS-lambda clone.

  Only relevant restriction sites are indicated that have been used for subcloning and sequencing of the MOX gene. The open reading frame, containing the structural DAS sequence, and the M13 subclones made, are depicted.
  - Fig. 20. Identical sequences in -1000 region of DAS and MOX genes.

#### CLAIMS

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- 1. Process for preparing an oxidoreductase by culturing a microorganism under suitable conditions, optionally concentrating the enzyme and collecting the concentrated enzyme in a manner known per se, characterized in that a microorganism is used that has been obtained by recombinant DNA technology, and which is capable of producing the oxidoreductase.
- 2. Process according to claim 1, characterized in that the microorganism is capable of producing at least one enzyme selected from the group consisting of
  - (1) alcohol oxidases,

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- (2) amine oxidases, including alkylamine oxidase and benzylamine oxidase,
- (3) amino acid oxidases, including D-alanine oxidase, lysine oxidase,
  - (4) cholesterol oxidase,
  - (5) uric acid oxidase,
  - (6) xanthine oxidase,
- 20 (7) chloroperoxidase, and
  - (8) aldehyde oxidase.
  - 3. Process according to claim 1 or 2, characterized in that the microorganism is a mould or yeast.
  - 4. Process according to claim 3, characterized in that a mould or yeast is selected from the group consisting of the genera <u>Aspergillus</u>, <u>Candida</u>, <u>Geotrichum</u>, Hansenula, Lenzites, Nadsonia, <u>Pichia</u>, Poria,
- 30 <u>Polyporus</u>, <u>Saccharomyces</u>, <u>Sporobolomyces</u>, <u>Torulopsis</u>, Trichospora and Zendera.
- 5. Process according to claim 4, characterized in that the mould or yeast is selected from the species

  Aspergillus japonicus, Aspergillus niger, Aspergillus oryzae, Candida boidinii, Hansenula anomala, Hansenula

polymorpha, Hansenula wingeii, Kloeckera sp. 2201 and Pichia pastoris.

6. Process according to any one of claims 1-5, characterized in that the microorganism is also capable of producing a dihydroxyacetone synthase enzyme, which promotes the formation of dihydroxyacetone from formaldehyde.

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- 7. Use of an oxidoreductase prepared by a process as claimed in any one of claims 1-5 in an oxidation process.
- 8. Bleaching composition including a fabricwashing detergent composition or hard-surface-cleaning
  composition having bleach activity, characterized in
  that it contains an oxidoreductase prepared by a
  process as claimed in any one of claims 1-5 and a
  substrate for that oxidoreductase.
  - 9. Microorganism, obtainable by recombinant DNA technology and being capable of producing an oxido-reductase suitable for use in a process as claimed in claims 1-5.
  - 10. Microorganism, obtainable by recombinant DNA technology and being capable of producing a dihydroxy-acetone synthase-enzyme suitable for use in a process according to claim 6, in addition to being capable of producing an oxidoreductase.
  - 11. Process for preparing a transformed microorganism as claimed in claim 9, characterized in that a
    DNA sequence coding for an oxidoreductase together with
    one or more other DNA sequences which regulate the
    expression of the structural gene is introduced into
    the microorganism via an episomal vector or integration

in the genome, such that the microorganism is capable of producing the oxidoreductase.

- 12. Process for preparing a transformed microorganism as claimed in claim 10, characterized in that
  a DNA coding for a dihydroxyacetone synthase-enzyme
  together with one or more other DNA sequences which
  regulate the expression of the structural gene is
  introduced into the microorganism via an episomal
  vector or integration in the genome, such that the
  microorganism is capable of producing the dihydroxyacetone synthase-enzyme (DHAS enzyme).
- 13. DNA sequence coding for an oxidoreductase, characterized in that it is obtainable by recombinant DNA technology from natural and/or cDNA and/or chemically synthesised DNA.
- 14. DNA sequence according to claim 13, charac-20 terized in that it codes for an alcohol oxidase.
- 15. DNA sequence according to claim 14, characterized in that it comprises the DNA sequence 1-1992 (MOX gene) given in Fig. 11A + 11B encoding the polypeptide 1-664 (MOX), the amino acid sequence of which is given in Fig. 11A + 11B.
- 16. Combination of DNA sequences comprising a structural gene coding for an oxidoreductase and one or more other DNA sequences which regulate the expression of the structural gene in a particular microorganism or group of microorganisms.
- 17. Combination of DNA sequences according to claim 16, characterized in that it comprises at least part of the upstream DNA sequence -1 to about -1500 given in Fig. 11A and/or at least part of the down-

stream DNA sequence 1993 to about 3260 given in Fig. 11B (regulatory regions of the MOX gene).

- 18. Combination of DNA sequences according to claim 17, characterized in that it comprises at least the polynucleotide -1052 to -987 of the upstream DNA sequence given in Fig. 11A.
- 19. Combination of DNA sequences according to
  10 claim 17, characterized in that it contains a modified
  MOX promoter sequence which is obtainable by deletion
  of at least polynucleotide -1052 to -987 given in Fig.
  11A.
- 15 20. Combination of DNA sequences according to claim 16, characterized in that it comprises at least part of the upstream DNA sequence -1 to about -2125 given in Fig. 18A + 18B and/or at least part of the downstream DNA sequence 2107 to about 2350 given in Fig. 18C (regulatory regions of the DAS gene).
  - 21. Combination of DNA sequences according to claim 20, characterized in that it comprises at least the polynucleotide -1076 to -937 of the upstream DNA sequence given in Fig. 18A.

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- 22. Combination of DNA sequences according to claim 20, characterized in that it contains a modified DAS promoter sequence which is obtainable by deletion of at least polynucleotide -1076 to -937 given in Fig. 18A.
- 23. Combination of DNA sequences according to claim 16, characterized in that it comprises a
  35 structural gene coding for an oxidoreductase of a higher eukaryote, a mould, or a yeast.

- 24. Combination of DNA sequences according to claim 23, characterized in that it comprises a structural gene coding for an oxidoreductase of a yeast of the genus <u>Hansenula</u>, preferably of the species <u>H</u>.
- 5 polymorpha.

- 25. Combination of DNA sequences according to claim 16, characterized in that the structural gene coding for an oxidoreductase encodes an alcohol oxidase.
- 26. Combination of DNA sequences according to claim 25, characterized in that the structural gene is the DNA sequence 1-1992 (MOX gene) given in Fig. 11A + 11B encoding the polypeptide 1-664 (MOX), the amino acid sequence of which is given in Fig. 11A + 11B.
- 27. Combination of DNA sequences according to claim 16, characterized in that it also contains a20 structural gene coding for DHAS.
- 28. Combination of DNA sequences according to claim 27, characterized in that it contains a structural gene coding for DHAS having the amino acid sequence as given in Fig. 18B + 18C.
- 29. Combination of DNA sequences according to any one of claims 16-28, characterized in that the DNA sequences have been modified, while retaining their coding function for an oxidoreductase or for their regulatory functions, by recombinant DNA technology.
- 30. Combination of DNA sequences according to any one of claims 16-29, characterized in that it contains one or more DNA sequences that enable stable inheritance of said combination in the progeny of any particular host organism.

- Combination of DNA sequences suitable for the transformation of a microbial host to produce a specific enzyme or other protein which combination of DNA sequences contains a regulon, a structural gene 5 coding for that specific enzyme or other protein and optionally a terminater, characterized in that a regulon is used selected from the group consisting of at least part of the regulon -1 to about -1500 of the MOX gene given in Fig. 11A or at least part of the 10 regulon of -1 to about -2125 of the DAS gene given in Fig. 18A and modifications thereof that do not impair the regulon function, and optionally a terminater is used selected from the group consisting of at least part of the terminater 1993 to about 3260 of the MOX 15 gene given in Fig. 11B or at least part of the terminater of 2110 to about 2350 of the DAS gene given in Fig. 18B and modifications thereof that do not impair the terminater function.
- 20 32. Combination of DNA sequences according to claim 31, characterized in that it is suitable for transformation of a <u>Hansenula</u> yeast, in particular a Hansenula polymorpha.
- 25 33. Combination of DNA sequences according to claim 31, characterized in that it is suitable for transformation of a Saccharomyces yeast, in particular Saccharomyces cerevisiae.
- 30 34. Combination of DNA sequences according to claim 31, characterized in that the structural gene coding for that specific enzyme or other protein contains DNA sequences derived from the structural gene coding for MOX (Fig. 11A + 11B), which modify said specific enzyme or other protein, without impairing its functions, in such a way that said specific enzyme or other protein is translocated into the peroxisomes or equivalent

microbodies of said microbial host.

- 35. DNA sequence coding for a dihydroxyacetone synthase-enzyme, characterized in that it is obtainable by recombinant DNA technology from natural and/or cDNA and/or chemically synthesised DNA.
- 36. DNA sequence according to claim 35, characterized in that it comprises the DNA sequence 1-2106

  (DAS gene) given in Fig. 18B + 18C encoding the polypeptide 1-702 (DHAS), the amino acid sequence which is given in Fig. 18B + 18C.
- 37. Combination of a DNA sequence coding for a dihydroxyacetone synthase-enzyme and one or more other DNA sequences which regulate the expression of the structural gene in a particular microorganism or group of microorganisms.
- 20 38. Combination of DNA sequences according to claim 37, characterized in that it comprises the DNA sequence according to claim 36 (DAS gene) and at least part of the upstream DNA sequence -1 to about -2125 given in Fig. 18A + 18B and/or at least part of the downstream DNA sequence 2107 to about 2350 given in Fig. 18C (regulatory regions of the DAS gene) and/or at least part of the upstream DNA sequence -1 to about -1500 given in Fig. 11A and/or at least part of the downstream DNA sequence 1993 to about 3260 given in Fig. 11B (regulatory regions of the MOX gene).
- 39. Combination of DNA sequences according to claim 38, characterized in that it comprises at least the polynucleotide -1076 to -937 of the upstream DNA sequence given in Fig. 18A or at least the polynucleotide -1052 to -987 of the upstream DNA sequence given in Fig. 11A, respectively.

- Process for preparing a polypeptide, such as a 40. protein or an enzyme, by culturing a microorganism under suitable conditions, optionally concentrating the polypeptide and collecting same in a manner known per se, characterized in that a microorganism is used that has been obtained by recombinant DNA technology and carries a structural gene coding for the polypeptide concerned, the expression of which is under the control of a regulon, comprising a promoter and at least either the -1052 to -987 region of the MOX gene 10 of Hansenula polymorpha CBS 4732, or the -1076 to -937 region of the DAS gene of Hansenula polymorpha CBS 4732, or a corresponding region of other methylotrophic moulds or yeasts, or an effective modification of any 15 of these regions.
  - 41. Process according to claim 40, characterized in that the promoter is derived from the yeast Hansenula polymorpha.

42. Process according to claim 40 or 41, characterized in that the microorganism is a mould or yeast.

43. Process according to any of claims 40-42,
25 characterized in that a mould or yeast is selected from
the group consisting of the genera Aspergillus,

Candida, Geotrichum, Hansenula, Lenzites, Nadsonia,
Pichia, Poria, Polyporus, Saccharomyces, Sporobolomyces, Torulopsis, Trichospora and Zendera.

44. Process according to claim 43, characterized in that the mould or yeast is selected from the species Aspergillus japonicus, Aspergillus niger, Aspergillus oryzae, Candida boidinii, Hansenula anomala, Hansenula polymorpha, Hansenula wingeii, Kloeckera sp. 2201 and Pichia pastoris.

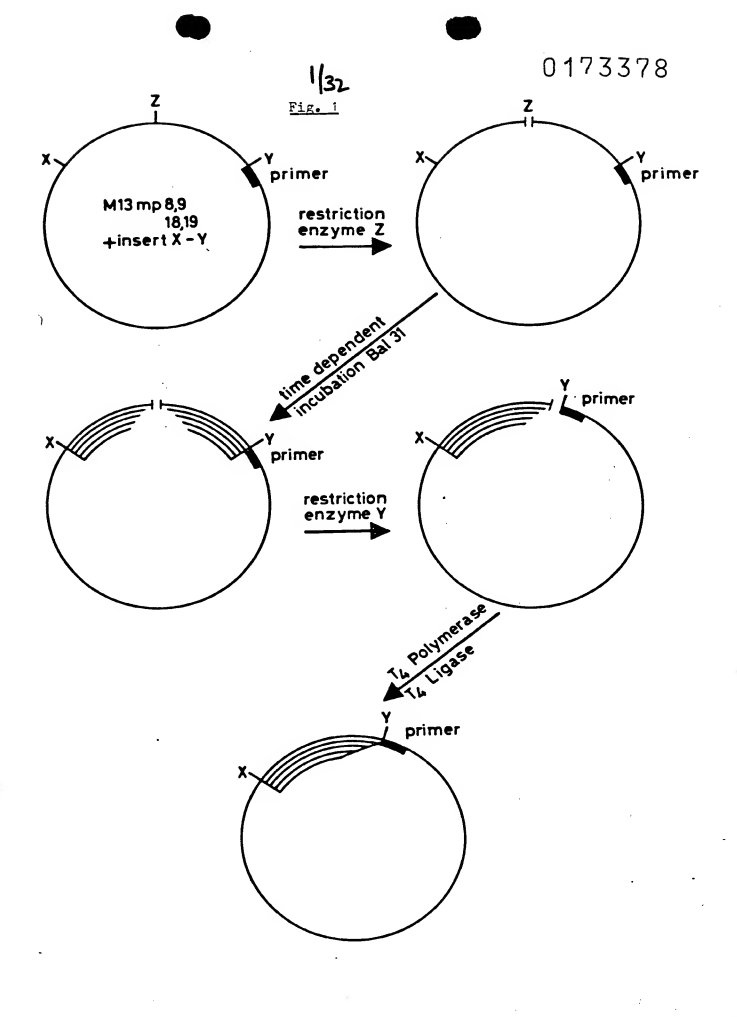
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- 45. Process according to claim 44, characterized in that the microorganism is the yeast species Hansenula polymorpha.
- 5 46. Process according to any of claims 40-45, characterized in that the structural gene concerned has been provided with one or more DNA sequences which translocate the gene product into the peroxisomes or equivalent microbodies of the microbial host.

47. Process according to claim 46, characterized in that the DNA sequences concerned consist of the MOX gene or those parts thereof which are responsible for MOX translocation into the peroxisomes or microbodies.

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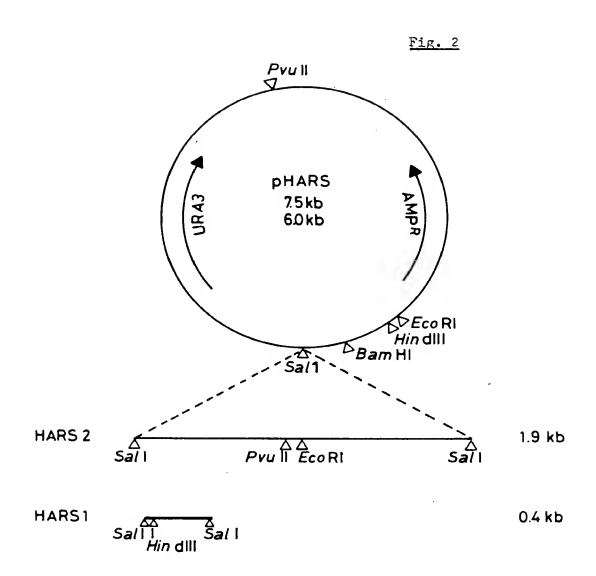
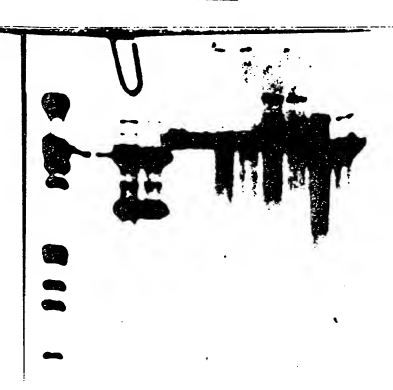


Fig. 3 DNA sequence of the autonomous-replicating sequence HARS1 from the methylotrophic yeast Hansenula polymorpha. The HARS1 represents a Sall fragment comprising 483 nucleotides. The dideoxy-sequencing method was employed.

(GTCGACTCCG CGACTCGGCG TTCACTTTCG AGCTATTCAT CAACGCCGGA ATACGTCAGA AACAGCCGTG CCCCAGGGAC CAGAAAGCCT ACTGGTGAGT ATGTTCTTTC GTGTGATTTT CCGAGGATGA GACGACGATA ACGAGCACAA CTCGGAGTCG GAGGACACGC TTATTGCGTT GACGAGCCAC ATCAGCAGGC TGTCAAGACT GAGTATAGGC CACAGAGCTG ATTCTGCTCA TACTCAAGAC GTTAGTAAAC TCCGTCTGCC ACAATGCTGA CAGAGTATTA TAATAATAGT GAATTACGAA CAATGTAGTC AAAAAAATTT AGTAACAATA TGTCATGATG ACAGATTTGC TGAAACCAGT GAACTCCAAT AAATCCAGCG GCTACCGCAT CCCAAGAGAA ACAGATCAGA GGTCTAGGCT TGTTTCAGAG TACTACAAGC TTTCCAGAAC TTAGCAATTC TCAAACGCGG TTTGCCGCAC

Fig. 4

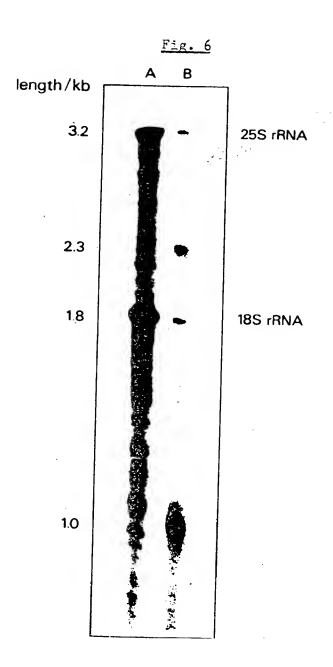


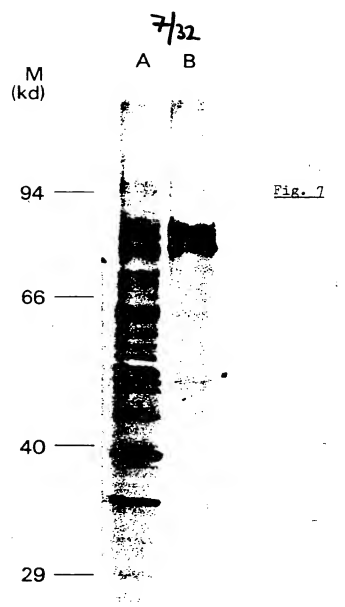
1 23 45 67 89 40 11 12 13

Fig. 5



1 2 3 4 5 6 7 8



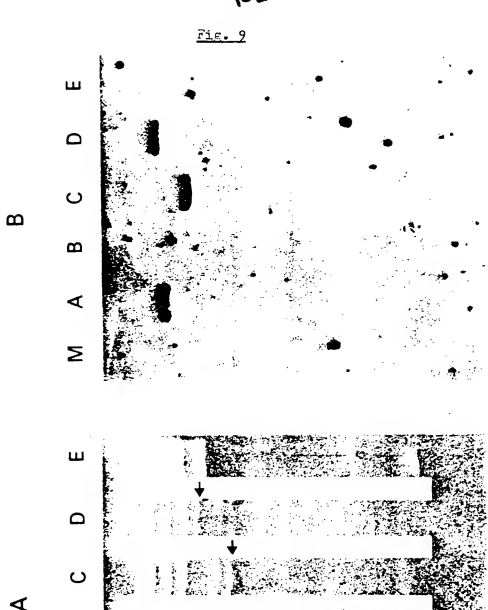


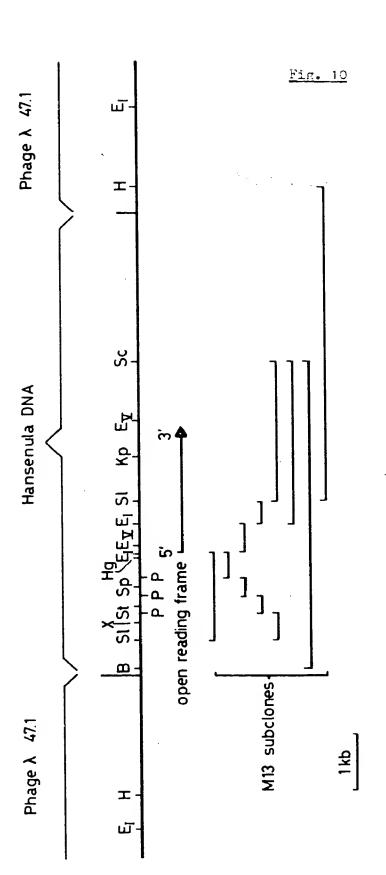
#### Fig. 8

NH2-Ala-Ile-Pro-Asp-Glu-Phe-Asp-Ile-Ile-Val-Val-Gly-

CCA GAC GAA TTC GA

-Gly-Gly- \* -Thr-Gly-Cys-Cys-Ile-Ala-Gly- \* -Leu--Ala-Asn-Leu-Asp-Asp-Gln-Asn-Leu





78

•	"	132	
		19 -	01733
- <u>- + K • 11A</u>	-1501	CTCCCGGATC AGCTTGTGGC	CCCGTAATCG
-1451		CACGCTCCCT CGCGAGCCCA	-1401
<b>-</b> ,		CAGTGTCAGA TGAATGTCCT -1351	
		TTTCCAGTAG AACCGTAATC	
•	-1251	CCCCCCCATA ATGAACACGT	
	-1201	GGCGTCGATG AGGCCCTTCA	
-1151		GTGACGATAC CCACATAGAG	-1101
		GACTCGGCAT CGAAATTTTT -1051	
	•	AGCTGCACCT TGCGATCTGG	
	-951	TGGGCCGTGT TGAGAATCAG	
	-901	CTGATGGCGC GGCGGTCGTC	
-851		AACCTGCCAA CGCTCGCAAC GTCATTTCAA TGTTGTCGTC	-801
-		-751 CGTTTTGCCT CAGAGTTTAC	
		-701 GCGTACAGGA CGAACGCCGT	
	-651	CGCTCTTCCT TGACTGCGTA	
	-601	CTGTGCTGGA TGCAGCAATT	•
-351		ACCCACTCGC TTTAGCCGTC	-501
		-451 GCCGCAACGG AACTTTTCCC	
		-401 TGCTGGGGTA GCCTACAACG	
	GCGCTACCCA CTCCTGTCTC	TGCCTCGTAC CAGAAAATCA	
ACAGGCCCGA	-301	CGGACAGCGT GTTTGTCCAC	
-251 CGGGTGAAC	GGCCAGAAAG TAAATTCTTA	TGCTAGCGTG CAGCGACTCC	-201 GACATCCCCA
GITTICCCC	TACTTCATCA CAGATGGGGT	-151 CAGCGCTGCC GCTAAGTGTA	CCCAACCGTC
CCCACACGG	CCATCTATAA ATACTGCTGG	-101 CACTCCAGGG TCCTGACATC	AATCTAAACT
1	-51 5	10	15
ACAAAAACAAA ATC GCC	ATT CCT GAG GAA TTC G	SP ILE ILE VAL VAL GLY AT ATC ATT GTT GTT GGT	GGA CGT TCC ACC
20	25 CIV ARC 150 ALA ARN 1	30 EU ASP ASP CLN ASN LEU	35
GCC TGC TGC ATT GCC	GGC AGA CTG GCA AAC C	TC GAC GAC CAA AAC CTC	ACA GIT CCC CIG
THE GLU GLY CLY GLU	ASM ASM ILE ASM ASM P	RO TRP VAL TYR LEU PRO	GLY VAL TYR PRO
ATC GAG GCT GCT GAG	AAG AAG ATC AAG AAC C	CT TGG GTC TAC CTT CCC	GGA CTG TAT CCT
ARG ASH MET ARC LEU	ASP SER LYS THE ALA T	HR PHE TYR SEN SER ARG	PRO SER LYS ALA
80	STAC TOO AND ACT TOO A	CC TTC TAC TCC TCC AGA	95

TRP SER THR ASP GLU LEU LEU PRO LEU ILE LYS LYS ILE GLU THR TYR CLW ARG PRO CYS
TGG AGC ACC GAC GAG TTG CTA CCT CTG ATC AAA AAA ATC GAA ACT TAC CAG CGT CCT TCC 140 145 150 155
ASH ASH ARG ASP LEU MIS CLY PHE ASP GLY PRO ILE LYS VAL SER PHE GLY ASH TYR THE
AAC AAC AGA GAT CTG CAC GCC TTT GAC GGC CCA ATC AAG GTT TCC TTT GGA AAC TAC ACG THE PRO THE CYS CLN ASP PHE LEU ARG ALA ALA GLU SEN GLN CLY ILE PRO VAL VAL ASP TAT CCT ACG TCC CAG GAC TTG CTG ACA GCA CAG TCG GAG GGA ATT CCT GTT GTG GAC 180 185 190 195 ASP LEU GLU ASP PHE LYS THE SEE WIS GLY ALA GLU MIS TRP LEU LYS TRP ILE ASH ARG GAC CTG GAG GAC TTC AAG ACA TCG CAT GGT GCA GAC GAC TGG CTG AAG TGG ATT AAC ACA 200 213 ASP LEU GLY ARG ARG SER ASP SER ALA HIS ALA TYR VAL HIS PRO THR HET ARC ASN LYS GAC CTC CGC AGA AGA TGG GAT TCT GCG GAC GCC TAC GTC CAC CCA ACT ATG ACA AAC AAG 220 225 230 235 CLH SER LEU PHE LEU ILE THR SER THR LYS CYS ASP LYS VAL ILE ILE GLU ASP GLY LYS
CAC AGC CTG TTC CTC ATC ACC TCG ACC AAG TGT GAC AAG GTG ATC ATC GAG GAC GGC AAG
240
ALA VAL ALA WAL ABC THR UAL TRA LYS CYS ASP LYS VAL ILE ILE GLU ASP GLY LYS
255 ALA VAL ALA VAL ARC THR VAL PRO HET LYS PRO LEU ASH PRO LYS LYS PRO VAL SER ARG GCT GTC GCC GTG AGA ACA GTG CCA ATG AAG CCT CTG AAC GCT AAG AAC CCT GTG TCC ACA 260 275 THE PME ARG ALA ARG LTS GLW ILE VAL ILE SER CYS GLY THE ILE SER SER PRO LEU VAL ACC TTC AGA GCC AGA AAG CAG ATT GTG ATC TCC TCC GGA ACC ATC TCG TCT CCT CTG GTG

280

280

280

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LEU GLH ARG SER GLY ILE GLY ALA ALA HIS HIS LEU ARG SER VAL GLT VAL LYS PRO ILE

CTC CAG ACA TCT GCT ATT GGT GCA GCT CAC CAC TTG AGA TCC GTC GGG GTC AAG CCA ATC

123378

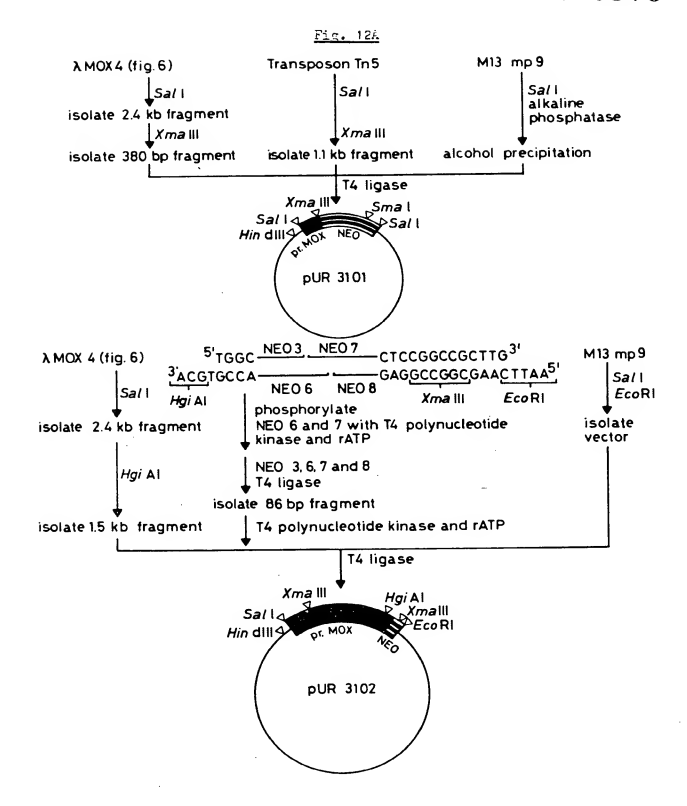
												•							
		300					305					310					315		()
					VAL														
CTC	CAC		GCA	CCT	CTC	CCI		AAT	TTC	CAG	GAC		TAC	TCT	TTC	TTC		CCA	TAC
		320					525					330					555		
					GTT														
120	010	340	001	UAL			343	110	CAC	CAC		350	ACC		CAC	CCA		CCC	CAG
LTS	ALA		PRE	ASP	CL M	TRP		222	ARR	1 7 8	ARD			1 # 11	T # B	T M B	355	CLT	11.
AAG	SCC	GCT	TTC	GAC	CAG	TCC	TAC	TCC	AAC	AAG	GAC	CET	CCA	TTG	ACC	ACC	AAC	CCT	ATT
		560					365					370					575		
CLD	ALA		VAL	LTS	ILE	ARG		TER	CLU	CLO	CLO		ALA	THR	ALA	ASP		ASP	PME
GAA	CCC	G CA	GTC	AAG	ATC	ACA	CCT	ACC	GAA	CAC	CAC	CTC	CCT	ACC	CCC	GAC	CAC	CAC	TTC
		380					385					390					395		
ARG	ARC	CLT	TTE	ALA	CLO	TTR	PHR	CLU	ASE	LTS	PRO	ASP	LTS	PRO	LEU	HET	RIS	TTR	SER
ACA	ccc	CCC	TAC	CCA	CAC	TAC	TTC	CAC	AAC	AAG	CCA	GAC	AAC	CCT	CTG	ATC	CAC	TAC	TCT
		AOO					405					410					415		
VAL	ILE	BER	GLT	POL	PHE	CLT	ABP	HIS	THE	LTS	ILE	PRO	ASH	CLT	LTS	PHE	MET	TRR	MET
CTC	ATC	TCC	GCC	TIC	TTT	CCA	GAC	CAC	ACC	AAC	ATT	CCT	AAC	CCC	AAC	TTC		ACC	ATC
		4 20		61.0	TTR		4 2 5		480	c		450					455		
776	545	776	CTC	CAC	TAT	CCA	TTC	700	ACA	CLI	PAL	VAL	ARG	ILE	IBE	BEK	ALA	ASH	PRO
110		440	010	CAC			445			CCA		450	AGA	AIC	ACC	106	455	AAC	CCA
TTR	ASP		PRO	ASP	PHE	ASP		CLT	PHE	LRD	ASH		61.0	ARG	ARP	1 7 13		-	***
					TTC														
		4 60					465		•••			470					475		***
VAL	TRP	ALA	TTR	LTS	LTS	SEE		CLU	TRR	ALA	ARC		MET	CLU		PRE		GLT	GL U
					AAG														
		ASO					485					490					495		
VAL	TRR	SER	H 14	RIS	PRO	LEU	PHE	LTS	VAL	ASP	SER	PRO	ALA	ARC	ALA	ARC	ASP	LEU	ASP
CTC	ACC	TCG	GAC	CAC	CCA	TTC	TTC	AAC	CTT	CAC	TCG	CCA	CCC	ACA	CCC	AGA	CAC	CTG	GAC
		500					503					510					515		
LEO	CLU	TRE	CTS	SER	ALA	TTR	ALA	CLT	PRO	LTS	HIS	LEU	THE	ALA	ASR	LEU	TTR	RIS	CLT
CTC	CAC	ACA	TCC	ACT	CCA	TAT	CCC	CCT	CCT	AAC	CAC	CTC	ACT	ccc	AAC	CTC	TAC	CAC	CGC
		520					525					5 50					555		
358	IRP	IBR	VAL	PRO	ILE	ASP	LTS	PRO	THE	PRO	LTS	ASR	ASP	PHE	RIS	VAL	THE	SER	ASM
1 00	100	340	OI I	CCI	ATC	CAC	AAG	CCA	ACG	CCT	AAC		GAT	TTC	CAC	CTC		TCC	AAC
GL B	WAI					4	343	e				550					555		
CAA	GTC	CAA	CTC	CAC	SER TCC	246	ATC	CLU	118	IBE	CLU	CLU	ABP	ASP	CLU	ALA	ILE	VAL	ASH
		560				CAL	363	CAC	1 4 5	ALL	CAC	370	GAC	CAC	CAG	ccc		CTC	AAC
TTR	ILE		GLU	RIS	THE	GLU		TRR	TEP		CTE	1 70	GIT	- 40	***		575		
TAC	ATT	AAG	GAA	CAC	ACC	GAC	ACG	ACT	TEE	CAC	TET	CTG	CCT	400	7.00	700	ATC	CCC	CCA
		380					585					590					595		
ARG	CLO	GLT	SER	LTS	ILR	ALA	PRO	LTS	CLT	CLT	VAL	LEU	ASP	ALA	ARC	LEU	ARR	VAL	TTR
ACA	CAC	CCT	ACT	AAC	ATT	CCT	CCT	AAC	CCA	CGT	CTC	TTC	GAC	GCC	AGA	CTG	AAC	CTT	TAC
		• 00					603					610					615		
CLT	VAL	CLE	ASM	LEU	LTS	VAL	ALA	ASP	TKU	SER	VAL	CTS	PRO	ASP	ASH	VAL	CLT	CTS-	ABM
CCA	CTC	CAG	AAC	CTC	AAG	CTT	CCC	GAC	CTT	TCT	CTT	TCT	CCC	CAC	AAC	CTT	CCA	TGC	AAC
		620				_	625					630					635		
TEE	TIE	SER	THE	ALA	LEU	THE	ILE	CLT	CLO	LTS	ALA	ALA	THE	LEU	VAL	ALA	CLU	ASP	LEU
ALL	IAC	ICI	ACT	CCA	TTC	ACC	ATC	CCT	CAG	AAC	CCT	CCC	ACT	CTT	CTT	CCT	CAA	CAT	CTT
CL Y	TTP	640	~1+				645					650					655		_
ccc	TAC	TCA	CCC	700	ASP	CIC	437	HEI	1 8 8	1 L L	PEO	ASE	PHE	ARC	LEO	CLY	THE	TTR	CLU
		660			CAC	-16	UAL	~ 1 G	466	AII	CCA	440	TIC	AGA	CTC	CCA	ACT	TAC	CAC
CLD	THE		LEU	ALA	ARG	PHF	•••											-	
GAG	ACC	GGA	CTT	GCC	AGA	TTC	TAA	GCAG	: AC	TCC	ACC	404	***		***				

GGA CTT GCC AGA TTC TAA GCAG ACGTGGAAGG ACATACCGCT TTTGAGAAGC 2000 CTCTTTCAAA ATACTTCTTT TTCTCCTTTA TATCCTTTAT CAACTCATCA CATCAAAACC 2050 TCAMATAGCG ACTATACGAM MATTTAMTGM AMATTMAMTT AMATMTTTC TTAGGCTATT AGTEACETTE AAAATGEEEG CEGETTETAA CAACGTTETE ATGATEGACA ACTACGACTE CTTTACCTGG AACCTCTACG AGTACCTGTC TCACCAGGGA CCCAATGTCG AGGTTTTCAG CAACGATCAC ATCACCATTC CGGAGATTGA CCAGCTGAAC CCGGACCTGA TCACCCATTT

CCCTGGTCCT CCCCATCCAA CAACAGACTC CCGAATATCT CGCGACCTGA TCACCCATTT

2350

TAAACGGAAC ATTCCTGTCT TTGGTGTCTC TATGGGCCAC CACTGTATCT TCGAGGAGTT TEGECGACAE CTEGAGTATE EGGGEGAGAT TETECATEGA AAAACETECA ETETTAAGEA COMPACAMENT GEARTETTEM AMARCETTEE CEMAGATETT GETETEMECA GATACEMETE 2550
CETEGECEGA ACGUTEMAGT EGETTECEGA ETEGETEMAGA ATEMETETE GEMEMBACAM 2600 CGGGATCATT ATGGGTGTGA CACACAAGAA GTACACCATC GAGGGCGTCC AGTTTCATCC 2650 AGAGAGCATT CTGACCGAGC ACCGCCATCT GATGATCCAG AATATCCTGA ACGTTTCCGG TEGTTACTEC GAGGAAAATE CCAACGCCC CCCTCAGAGA AACGAAAGCA TATTGGAGAA AATATACGGG CAGAGACGAA AAGACTACGA CTTTGAGATG AACAGACCGG GGCGCAGATT TGCTGATGTA GAACTGTACT TGTCCATGGG ACTGCACCGC CCCTAATGAA TTTTTACGAC AGATTGGAGC AGAACATCAG CGCCGCCAAG GTTGCAATTC TCAGCGAAAT CAACAGAGGG TEGECETETA AAGGEGTEAT EGAEGGAGAE GETAACGETG CEAAACAGGE CETCAACTAC GCCAAGGCTG GAGTTGCCAC AATTTCTCTT TTCACCGAGC CAACCTGGTT TAAACGAAAT ATCCAGGACC TEGAGETECC CACAAAAGGC ATTCACTCTG TECECAATAC ACCETGTATT 3150 TIGGGGAAGG ACTITATETT CAACAAGTAC CAAATTETAG ACCCCGACT GGCCCGACCA GACACGGTTC TGCTGATTGT CAACATGCTG AGCTC





#### Fig. 12B

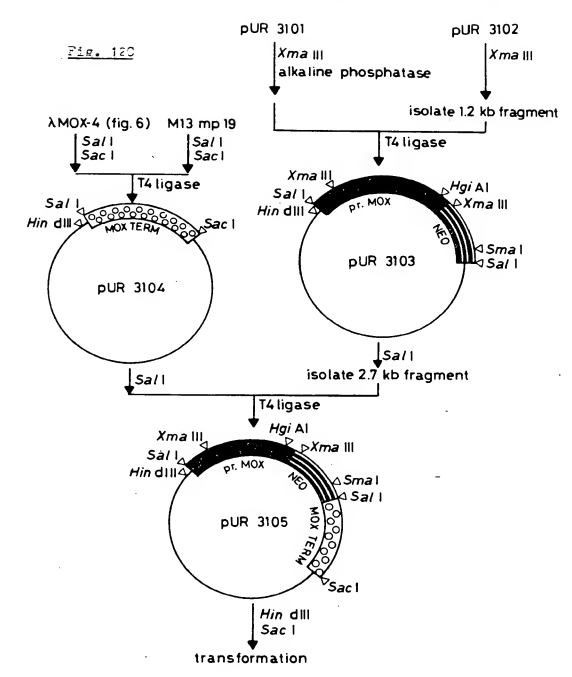
Promoter MOX-Neomycinphosphotransferase adaptor fragments

NEO3 5'CGGTGGTGACATCAATCTAAAGTACAAA 3'

NEO6 5'TCATTTTGTTTTTGTACTTTAGATTGATGTCACCACCGTGCA 3'

NEO7 5'AACAAAATGATTGAACAAGATGGATTGCACGCAGGTTCTCCGGCCGCTTG 3'

NEO8 5'AATTCAAGCGGCCGGAGAACCTGCGTGCAATCCATCTTGTTCAA 3'



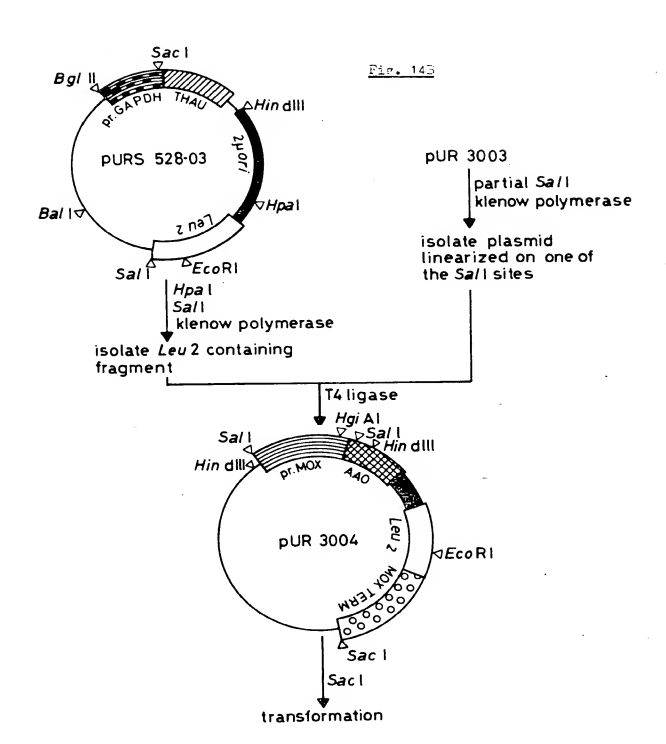
#### Pic. 13

			_		
< -34	PRO	HOTER MOX/A	AO ADAPTOR		>>
CGGTG	G TGACATCAA C ACTGTAGTT	T CTAAAGTAC	1 A AAAACAAAAT T TTTTGTTT <u>TA</u>	GAGAGTTGT	GTTATTGGTG
Hgial		- chillonio	He	t CICICAACAC	CAATAACCAC
<<	>				
CCGGTGTCA	T CGGTCTGTC	ACCGCCCTG	62 GTATCCACGA	GAGATACCAC	тссеттстес
GGCCACAGT	A GCCAGACAGO	IGGCGGGACA	CATAGGTGCT	CTCTATGGT	AGGCAAGACG
			122		
AGCCTCTGG	CGTTAAGGT	TACGCCGAC	CATTCACCCC	TTTC . CC . CC	ACCGACGTTG
I COUNCACE.	I GUARITUCA	ATGCGGCTGT	CTAAGTGGGG	AAAGTGGTGG	TGGCTGCAAC
CCGCCGGTC	T GTGGCAGCC	TACACCTCC	182 AGCCTTCCAA	CCCTCACCAC	
GGCGGCCAGA	CACCGTCGG	ATGTGGAGG	TCGGAAGGTT	GGGAGTCCTC	CGGTTGACCT
*******			242		
TGGTCGTCTC	GAAGTIGAT	CTCCTCTCC GAGGAGAGG	ACATCGGTTC TGTAGCCAAG	GCCTAACGCC	GCCAACATGG
				000N110000	COGITGIACC
GTCTGACCC	TGTCTCGGGT	TACAACCTGT	302 TCAGAGAGGC	CGTTCCTGAC	CCTTACTGGA
CAGACIGGG	ACAGAGCCCA	ATGTTGGACA	AGTCTCTCCG	GCAAGGACTG	GGAATGACCT
AGGACATGGT	CCTCGGTTTC	AGAAACCTTA	362 CCCCTAGAGA	CCTCC+C+T0	######################################
TCCTGTACCA	GGAGCCAAAG	TCTTTCGAAT	GGGGATCTCT	CGACCTGTAC	AAGGGACTGA
		Hindill			
ACAGATACGG	TIGGTTCAAC	ACCTCCCTGA	422 TCCTGGAGGG	TACA AACTAC	CTCC+CTCCC
TGTCTATGCC	AACCAAGTTG	TGGAGGGACT	AGGACCTCCC	ATCTTTCATG	GACGTCACCG
20.000			482		
ACTGGCTCTC	ACTGACCGAG TGACTGGCTC	AGAGGTGTTA TCTCCACAAT	AGTTCTTCCT TCAAGAAGGA	GAGAAAGGTC	GAGTCCTTCG
				O TO TITLONG	CICAGGRAGC
AGGAGGTTGC	CAGAGGTGGT	GCCGACGTCA	542 TCATCATGTG	TACCGGTGTC	TGGGCCGGTG
TOCTCOARCG	GICICCACCA	CGGCTGCAGT	AGTAGTACAC	ATGGCCACAG	ACCCGGCCAC
TCCTGCAGCC	TGACCCTCTG	. CTGCAGCCCG	602 GGAGAGGTCA	C A T C A T T A A C	
AGGACGTCGG	ACTGGGAGAC	GACGTCGGGC	CCTCTCCAGT	CTAGTAATTC	CAACTGCGGG
		XmaI			
CATGGCTGAA	GAACTTCATC	ATTACCCACG	ACCTGGAGAG	AGGTATCTAC	AACTCCCCTT
GTACCGACTT	CTTGAAGTAG	TAATGGGTGC	TGGACCTCTC	TCCATAGATG	TTGAGGGGAA
ACATTATCCC	TCCTCTCCAC		722		
TGTAATAGGG	ACCAGACGTC	CGGCAGTGGG	TGGGTGGTAC ACCCACCATG	CTTCCAGGTC GAAGGTCCAG	GGTAACTGGA CCATTGACCT
			KpnI	_	
ACGAGATCAA	CAACATCCAG	CACCACAACA	782 CCATCTGGGA		
TGCTCTAGTT	GTTGTAGGTC	CTGGTGTTGT	GGTAGACCCT	GGGTTGTTGT CCCAACAACA	AGACTGGAGC TCTGACCTCG
			842		
GATGGGACTT	GGACGCCAAG CCTGCGGTTC	ATCGTTGGTG TAGCAACCAC	AGTACACCGG TCATGTGGCC	TTTCAGACCT	GTTAGACCTC
				NANG TO TO GA	CARICIGGAG
AGGTCAGACT	GGAGAGAGA	CAGCTGAGAT	TCGGTTCCTC	CAACACCGAG	GTCATTCACA
LUCAGICIGA	CCTCTCTCTC	GTCGACTCTA	AGCCAAGGAG	STIGIGGCIC	CAGTAAGTGT
ACTACGGTCA	CGGTGGTTAC	GGTCTGACCA	962 TCCACTTGGG	FTCTC000==	0.000000
TGATGCCAGT	GCCACCAATG	CCAGACTGGT	AGGTGAACCC	ACACGGGAC	CTCCAACGGT
			1022		
AGCTGTTCGG TCGACAAGCC	TAAGGTCCTG ATTCCAGGAC	GAGGAGAGAA CTCCTCTCTT	ACCTGCTGAC (	CATGCCTCCA	TCCCACCTGT
				nosunuu f	AGGGIGGACA
GAG CTCAGCT					

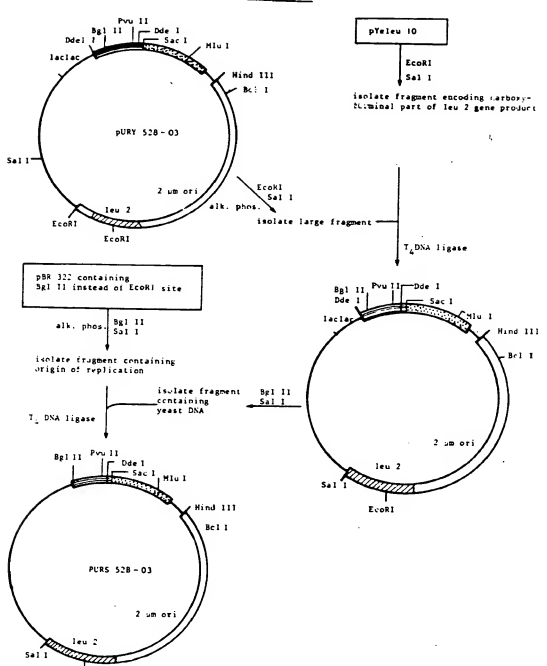
GAG CTCAGCT

Fig. 14A Xma| Kpn| Sall Hind III4 pUR 3001 pUR 3104 Sall Sal I isolate 1 kb fragment Hin dlll Sal I T4 ligase adaptor pr MOX-AAO Hindlig CAGCT5 T4 ligase Sac I pUR 3002 isolate 73 bp fragment pUR 3102 Sal I partial Sal 1 Hgi Al isolate 1.4 kb fragment containing promoter isolate linearized plasmid T4 ligase Hgi Al Sall Sall Hin dll Hin dlll4 pUR 3003 Saci Sacl transformation









EcoRI

<---->
-34

CGGTG GTGACATCAA TCTAAAGTA CAAAAACAAA
ACGTGCCAC CACTGTAGTT AGATTTCAT GTTTTGTTT
Hg1AI

ATGTACGCCG ACGCCATCTT CACCAACTCC TACAGAAAGG TTCTGGGTCA GCTCTCGGCC TACATGCGGC TGCGTAGAA GTGGTTGAGG ATGTCTTTCC AAGACCCAGT CGAGAGCCGG

6.1

TCTITCGAAG ACGICCTGTA GTACAGCTCT GTCGTCCCAC TCAGGTTGGT CCTCTCCCA Hindiii Pati AGAAAGCIIC IGCAGGACAI CAIGICGAGA CAGCAGGGIG AGICCAACCA GGAGAGAGGI

121

GCCAGAGCCA GACTGTGAG CGGTCTCGGT CTGACACTCA GCT

\*\*\* Sall

ig. 15

Fig. 16A

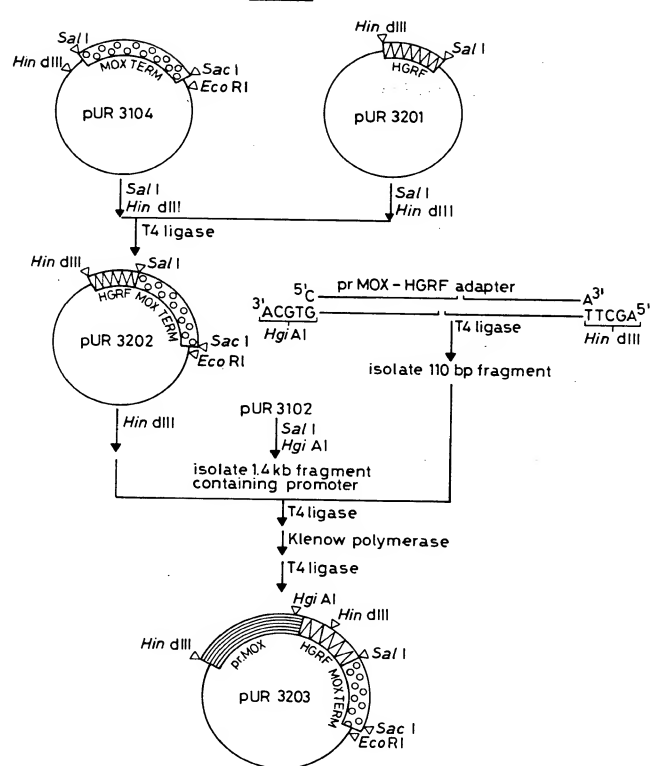
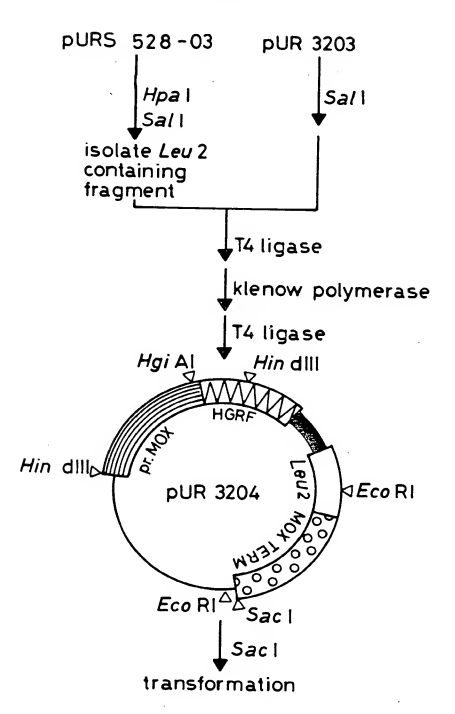


Fig. 16B



#### Fig. 160

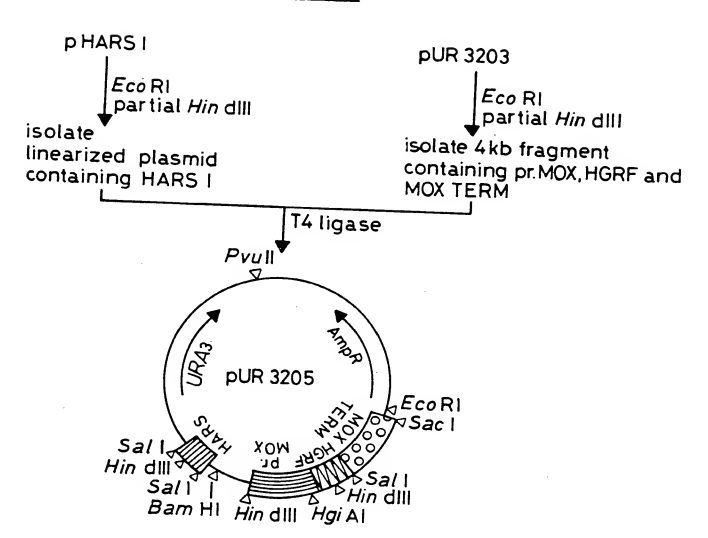
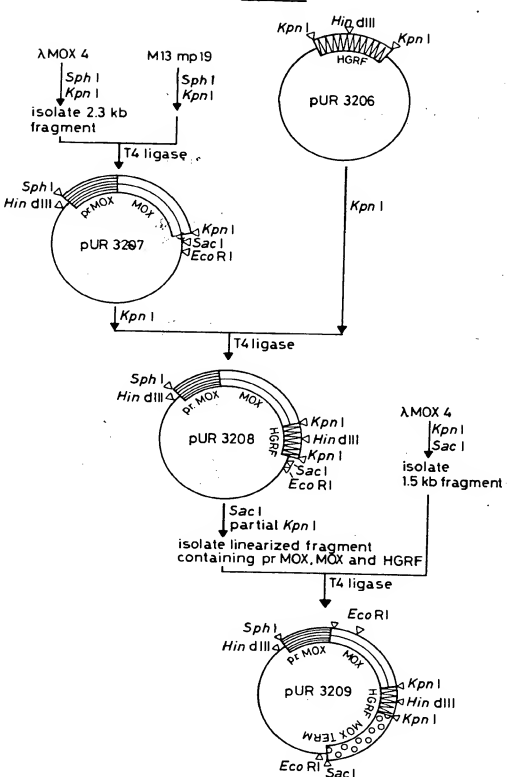
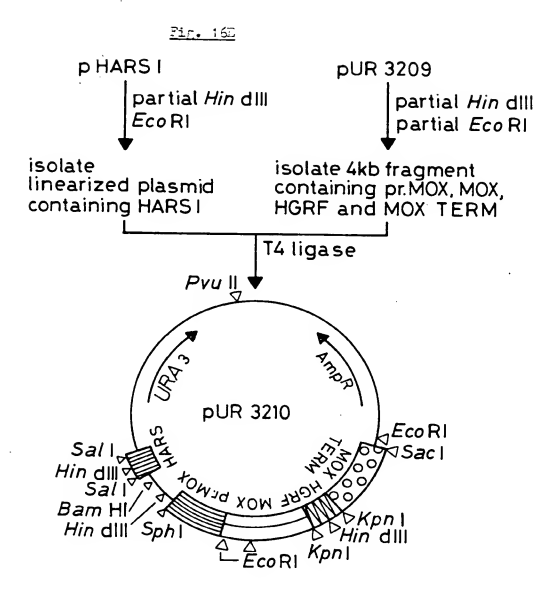
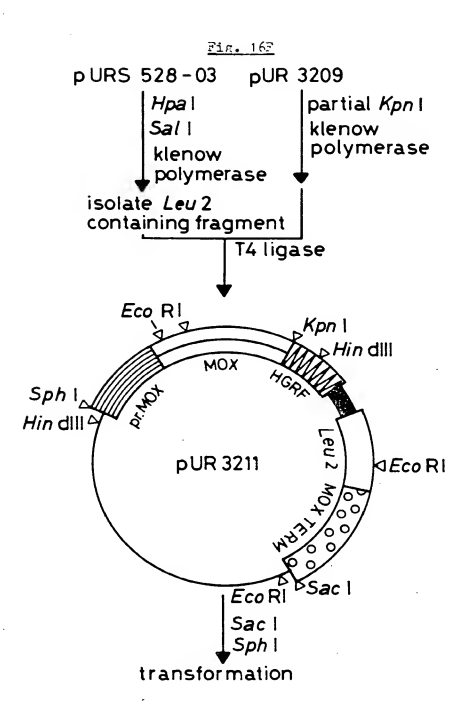


Fig. 16D







CATGTACGCCG ACGCCATCTT CACCAACTCC TACAGAAAGG TTCTGGGTCA GCTCTCGGCC CATGGTACATGCGCC TGCGGTAGAA GTGGTTGAGG ATGTCTTTCC AAGACCCAGT CGAGAGCCGC

AGAAAGCTTC TGCAGGACAT CTGTTCGAGA CAGCAGGGTG AGTCCAACCA GGAGAGGT TCTTTCGAAG ACGTCCTGTA GACAAGCTCT GTCGTCCCAC TCAGGTTGGT CCTCTCCCA Hindiii Pati

GCCAGAGCCA GACTGTGAGGTAC CGGTCTCGGT CTGACACTC \*\*\* KpnI

# 28/32

	Fig.	18A			
					GATCCACCTG
CTTGGCCAAT	GATTCAGCTG	CTGGACCGAA	AACGCCTCTT	TTGGCCAAAA	2125 AAAGCCCACC
GTTGATAACT -2054	GCGGAGGCCA	TATTTCAAAG	AACAGCGAAT	AAGAAAAAA	GGTGAATGAA
ATGCGCGAAA	CGATACCACT	TATTAGCATA	AACAAAAAA	AAAAAAATCT	ATTAGCTGTT
	GTTCAATAAT		-1904	GCGGCCTATT	GTCATCAGTG
	AACAGGTAAA	-1854			TTCCCAGACA
	TTCCGTGTCC -1804	• •			
-1/34	CTTCACCACG				-1704
	CCAGTTCTCC			-1654	GTGCAAGACG
	GATGAGAGA		-1604		
	CAACCTGGGA	-1554			
	GCTCGAGAGC -1504				
-1434	TGCGGAGATG				-1404
	GAGAAGGCGT			-1354	
	AATTTTTAAT		-1304		
GCGTCAAGCT		-1254	ACAGTCATGG		
	TGGCATATAT				
-1154	CCACCAGTGA				-1104
	TGTCCGTGTA			-1054	
	CCCTCTTCTC		-1004		
	TCGCTTCGTA	-954			
	TTTTCCGTCC -904				
-854	TAATCGGCTC			•	-804
	TGATGTCTTC			-754	
	GCTCTTCTGG		-704		
	CTCCGTAAAA	-654			
	AAATTGCCCA -604				
-554	CGGAAATTCC				-504
	CGTAGTATCA			-454	
	TCGAGGAAAG		-404		
	ACGCTGGCGT	-354			
	CCACCACCCT -304				
-254	ATCTGAAACC	GATGAAACGG	ACGACACTCC	CAACAAGCTC	ACTGCACTAT

### Fig. 183

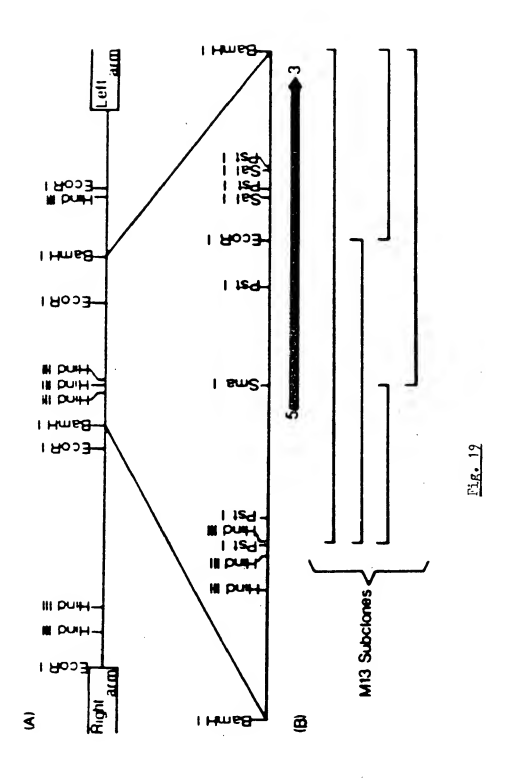
TTTTTTTTC TAGTGAAATA GCCTATCCTC GTCTGCTCC CCTCATACCT GTAAAGGGGT -154
GCAATTTAGC CTCGTTCCAG CCATTCACGG GCCACTCAAC AACACGTCGG CTACCATGGG
GTGCTTGGGC ACCAAAAGGC CTATAAATAG GCCCCCATCC GTCTGCTACA CAGTCATCTC

							•	-54				CALC		CIGC	IACA	CAG	ICAI	CIC	
				1				5					10					1.5	
				HET	SER	MET	ARG	ILE	PRO	LYS	ALA	ALA	SER	VAL	ASN	ASP	GLU	61.5	RIS
IG	ICII	TTCT	TCCC	ATG	ACT	ATG	AGA	ATC	CCT		GCA	GCG	TCG	GTC	AAC	GAC	GAA	CAA	CAC
-14	•		20																
C1 H	ARC	T1 P						25					30					35	
CAG	AGA	ATC	ILE	AAG	TAC	GLI	ARG	ALA	LEU	VAL	LEU	ASP	ILE	AVF	GLU	GLN	TYR	GLY	GLY
			ATC	, and	140	001	CGI	45	CII	CIC	CIG	GAC	ATT	GIC	GAG	CAG	TAC	GGA	GGA
GLY	BIS	PRO	GLY		A1. A	MPT	GI ¥	41.4	W				50					5 5	
GGC	CAC	CCG	GGC	TCG	GCC	ATG	CCC	CCC	ATC	CCT	ATC	GLY	ILE	ALA	LEU	TRP	LYS	TYR	THR
								0.0					70						ACC
LEU	LYS	TYR	ALA	PRO	ASN	ASP	PRO	ASW	TYR	PRE	ASR	ARG	ACB	ARG	985	WAT	1 22 21	75	
CTG	AAA	TAT	001	CCC	AAC	GAC	CCT	AAC	TAC	TTC	AAC	AGA	GAC	AGG	TTT	GTC	CTG	SER	
			00					83					80						
GLY	HIS	VAL	CYS	LEU	PHE	GLN	TYR	ILE	PHE	CLN	HIS	LEU	TYR	GLY	LEU	LYS	SER	MET	THR
661	CAC	GIG		CIG	TTC	CAG	TAT	AIC	TTC	CAG	CAC	CIG	TAC	GGT	CTC	AAG	TĆG	ATG	
			100					105					110					115	
ATG	GCG	CAG	LEU	AAG	TCC	TAC	HIS	SER	ASN	ASP	PHE	HIS	SER	LEU	CYS	PRO	CLY	HIS	PRO
		0.00	CTG 120	AAU	100	IAC	CAC	125	AAT	GAC	TIC	CAC	TCG	CTG	TGT	ccc	CCT	CAC	CCA
GLU	ILE	GLU	HIS	ASP	ALA	VAT.	GI II	VAI	700	<b>T D D</b>	C: ¥		130					135	
GAA	ATC	GAG	CAC	GAC	GCC	GTC	GAG	GTC	ACA	ACC	CCC	FRU	CTC	GLY	GLN	GLY	ILE	SER	ASN
								193					1 5 0						
SER	VAL	CLY	LEU	ALA	ILE	ALA	THR	LYS	ASR	LEU	ALA	A1.A	TDD	TYP	ACD	1 46	B B A	155	
TCT	GII	GGT	CIG	CCC	ATA	CCC	ACC	AAA	AAC	CTG	GCT	GCC	ACG	TAC	AAC	AAG	CCG	GEC	TTT
			100					100					170						
ASP	ILE	ILE	THR	ASR	LYS	AVF	TYR	CYS	MET	VAL	GLY	ASP	ALA	CTS	LEU	GLN	GLU		PRO
GAT	ATC	ATC	700	AAC	AAG	CTG	TAC	166	ATG	CII	CCC	GAT	CCG	TGC	TTG	CAG	GAG	GGC	CCT
			100					122					100						
GCT	CTC	GAG	SER	ATC	TCC	LEU	ALA	GLY	HIS	MET	GLY	LEU	ASP	ASN	LEU	ILE	VAL	LEU	TYR
	0.0	UNU	TCG 200	AIC	100	C16		205	CAC	ATG	CCC	CTG	GAC	AAT	CTG	ATT	GTG		TAC
ASP	ASN	ASN	GLN	VAL	CYS	CYS	ASP	CIV	CFD	<b>W</b> A 7	465		210					215	
GAC	AAC	AAC	CAG	GTC	TGC	TGT	GAC	GGC.	AGT	GTT	VOL	ATT	ALA	ASN	THR	GLU	ASP	ILE	SER
								443					710					336	ACT
ALA	LYS	PHE	LYS	ALA	CYS	ASN	TRP	ASH	VAL.	ILE	GLII	WAT	CIN	A C 10	41.4		A1 15	235	
CCC	AAG	TTC	AAG	CCC	TGC	AAC	TGG	AAC	GTG	ATC	GAG	GTC				TCC			VAL GTG
			240					245					250					255	
ALA	THR	ILE	VAL	LYS	ALA	LEU	CLU	TYR	ALA	CLN	ALA	CLU	LYS	HIS	ARG	PRO	TRR		TIP
OC1	ACC	ATT	GTC 260	AAG	CCC	TIG	GAG	INC	CCC	CAG	CCC	CAG	AAG	CAC	AGA	CCA	ACA	CII	ATC
AAC	TGC	AGA	THR	GTG	ATT	CCA	SER	GLY	ALA	ALA	PHE	CLU	ASN	HIS	CYS	ALA	ALA	HIS	GLY
			ACT 280			UUA	100	285	GCI	GCG	IIC	GAG	AAC	CAC	TGT	GCT	CCC		CCT
ASH	ALA	LEU	GLY	SLU	ASP	GLY	VAI	ARG	GIR	7 17 11			290					295	-
AAC	GCT	CTG	GGC	GAG	GAC	GGT	GTG	CGC	GAC	CTC	LIS	ATC	LYS	TYR	GLY	HET	ASH	PRO	ALA
												W I C	310	TAC	GGC	ATG	AAC		CCC
GLN	LYS	PHE	TYR	ILE	PRO	GLN	ASP	VAL	TYR	ASP	PHE	PHE	1 Ve	CI II	1 46	PRO		315	
CAC	AAG	TTC	TAC	ATT	CCG	CAG	GAC	GTG	TAC	GAC	TTC	TTC	AAG	GAG	AAG	CCC	CCC	CAC	CCC
ASP	LA2	LEU	VAL	ALA	CLU	TRP	LYS	SER	LEU	VAL	ALA	LYS		VAL	LYS	ALA	TYR		G 1. II
UAC	AAG	CIG		CCC	GAA	TCC	AAG	VO I	CTC	GTG	CCC	AAG	TAC	GTC	AAG	GCG	TAC	CCT	GAG
								367					110						
GAG	CCC	CAC	GLU	FHE	LEU	ALA	ARG	MET	ARG	GLY	CLU	LEU	PRO	LYS	ASN	TRP	LYS		PRE
- 70		CAG	GAG 360	III	ITC	CCC	CGC	AIG	AGA	CCC	GAG	CTG	CCV	AAG	AAC	TGG	AAG	TCG	TTC
			300					365					370					375	

#### Fig. 180

```
LEU PRO GLN GLU PHE THR GLY ASP ALA PRO THR ARG ALA ALA ARG GLU LEU VAL
CTG CCG CAG CAG GAA TTC ACC GGC GAC GCT CCT ACA AGG GCC GCT GCC AGA GAG CTT GTG
           380
                                385
                                                     390
                                                                         395
ARG ALA LEU GLY GLN ASN CYS LYS SER VAL ILE ALA GLY CYS ALA ASP LEU SER VAL SER
AGA GCC CIG GGG CAG AAC IGC AAG ICG GIG ATT GCC GGT IGC GCA GAC CIG ICI GTG
            400
                                405
                                                     410
                                                                         415
VAL ASN LEU GLN TRP PRO GLY VAL LYS TYR PHE MET ASP PRO SER LEU SER THR GLN CYS
GTC AAT TIG CAG TGG CCA GGG GTG AAA TAT TTC ATG GAC CCC TCG CTG TCC ACG CAG TGT
            420
                                425
                                                     430
                                                                         435
GLY LEU SER GLY ASP TYR SER GLY ARG TYR ILE GLU TYR GLY ILE ARG GLU HIS ALA MET
GGC CTG AGC GGC GAC TAC TCC GGC AGA TAC ATT GAG TAC GGA ATC AGA GAA CAC GCC
            440
                                445
                                                     450
CYS ALA ILE ALA ASN GLY LEU ALA ALA TYR ASN LYS GLY THR PHE LEU PRO ILE THR SER
TGT GCT ATC GCC AAT GGC CTT GCC GCC TAC AAC. AAG GGC ACG TTC CTG CCG ATC ACG
            460
                                465
                                                     470
                                                                         475
THR PHE PHE MET PHE TYR LEU TYR ALA ALA PRO ALA ILE ARG MET ALA GLY LEU GLN GLU
ACT TIC TIC ATG TIC TAC CTG TAC GCT GCC CCA GCC ATC AGA ATG GCC GGC CTG CAG GAG
            480
                                485
                                                     490
                                                                         495
LEU LYS ALA ILE HIS ILE GLY THR HIS ASP SER ILE ASN GLU GLY GLU ASN GLY PRO THR
CTC AAG GCG ATC CAC ATC GGC ACC CAC GAC TCG ATC AAT GAG GGT GAG AAC GGC CCT ACG
            500
                                505
                                                    510
                                                                         515
HIS GLN PRO VAL GLU SER PRO ALA LEU PHE ARG ALA TYR ALA ASN ILE TYR TYR MET ARG
CAC CAG CCG GTC GAG TCG CCA GCA TTG TTC CGG GCC TAT GCA AAC ATT TAC TAC ATG AGA
            520
                                525
                                                     530
                                                                         535
PRO VAL ASP SER ALA GLU VAL PRE GLY LEU PHE GLN LYS ALA VAL GLU LEU PRO PHE SER
CCG GTC GAC TOT GCA GAA GTG TIT GGC CTG TTC CAA AAA GCC GTC GAG CTG CCA TTC
            540
                                545
                                                    550
SER ILE LEU SER LEU SER ARG ASN GLU VAL LEU GLN TYR LEU ALA SER ARG ALA GLN ARG
TCG ATT CTG TCG CTC TCG AGA AAC GAG GTG CTG CAA TAC CTG GCA AGT CGA GCG CAG
            560
                                565
                                                     570
                                                                         575
ARG ARG ASN ALA ALA GLY TYR ILE LEU GLU ASP ALA GLU ASN ALA GLU VAL GLN ILE ILE
AGG CGC AAC GCG GCC GGC TAT ATT CTG GAG GAT GCG GAG AAC GCC GAG GTG CAG ATT ATT
            580
                                585
                                                     590
GLY VAL GLY ALA GLU MET GLU PHE ALA ASP LYS ALA ALA LYS ILE LEU GLY ARG LYS PHE
GGA GTT GGT GCA GAG ATG GAG TTT GCA GAC AAG GCC GCC AAG ATC TTG GGC AGA AAG TTC
            600
                                605
                                                     610
                                                                         615
ARG THR ARG VAL LEU SER ILE PRO CYS THR ARG LEU PHE ASP GLU GLN SER ILE GLY TYR
AGG ACC AGA GIT CTC TCC ATC CCA TGC ACG CGG CTG TIT GAC GAG CAG TCG ATC GGC TAT
            620
                                625
                                                     630
                                                                         635
ARG ARG SER VAL LEU ARG LYS ASP GLY ARG GLN VAL PRO THR VAL VAL VAL ASP GLY HIS
AGA CGC TCG GTT TTG AGA AAG GAC GGC AGA CAG GTG CCA ACG GTG GTG GTG GAC GGC CAC
            640
                                645
                                                     650
VAL ALA PHE GLY TRP GLU ARG TYR ALA THR ALA SER TYR CYS MET ASN THR TYR GLY LYS
GTT GCG TTC GGC TGG GAG AGA TAC GCT ACG GCG TCC TAC TGT ATG AAC ACG TAC GGC AAG
           660
                                665
                                                     670
                                                                         675
SER LEU PRO PRO GLU VAL ILE TYR GLU TYR PHE GLY TYR ASN PRO ALA THR ILE ALA LYS
TCT CTG CCT CCA GAA GTG ATC TAC GAG TAC TIT GGA TAC AAC CCG GCA ACG ATT GCC AAG
            680
                                685
                                                     690
                                                                         695
LYS VAL GLU ALA TYR VAL ARG ALA CYS GLN ARG ASP PRO LEU LEU HES ARG LEU PRO
AAG GTC GAA GCG TAC GTC CGG GCG TGC CAA AGA GAC CCT TTG CTG CTC CAC CGA CTT CCT
            700
GLY PRO GLU GLY LYS ALA ***
GGA CCT GAA GGA AAA GCC TAA CCACGAT AAAGTAAATA AGCTCTGATT AAGTAAGATG
                           2110
```

AATAAGTTET TIGIETGTGA ATGEEACECE ACAATAACEC CACAAATAAA ACTITEACAC TIGCGTCAGA AACTGTCGAG CCGCACGGGA CTGACTGTTT GGCGGCGTGC CTCTGTCCCC ACACGGATAT TICGCACGGA ACAGAAACCA TIGGACAAGG GGTTGCTGCC GATACCAAAT AGAATGCATC GGATCC



#### Fig. 20

### Identical sequences in -1000 region of DAS and MOX genes

DAS -937 ATCGCTTCGTACTCGCTCTGCAGCTTCGA

\*\*\*\* \*\*\* \*\* \*\*\*\*\*\* \*\*\*
ATCGAATGTAATGAGCTGCAGCTTGCGA

MOX -987

 $\mathbf{Application} \, \mathbf{0.1733378}$ 

85201235.0

# DECLARATION PURSUANT TO RULE 28, PARAGRAPH 4, OF THE EUROPEAN PATENT CONVENTION

The applicant has informed the European Patent Office that, until the publication of the mention of the grant of the European patent or until the date on which the application has been refused or withdrawn or is deemed to be withdrawn, the availability of the micro-organism(s) identified below, referred to in paragraph 3 of Rule 28 of the European Patent Convention, shall be effected only by the issue of a sample to an expert.

#### IDENTIFICATION OF THE MICRO-ORGANISMS

Accession numbers of the deposits:

CBS 7171

CBS 7472

AT CC 34438





0173378

## Unilever N.V.

Patent Division PO Box 137 3130 AC Vlaardingen The Netherlands

Olivier van Noortlaan 120 Tel. 010-60 69 33, (UTN Code 322) Telex 23261, Fax (gr II) 605800 Cables: Unileverpatent



European Patent Office P.O. Box 5818 2280 HV Rijswijk PLEASE ENDORSE AND RETURN DUPLICATE COPY AS ACKNOWLEDGEMENT.



Your Ref

Our Ref JVT/PS

Direct Dial. 60 . . . 5741Date

29 Aug. 1985

Re.: Recently filed patent application No. 85201235.0 - our case T 7000 (R)-EP

In this patent specification a Fig. 3 was present giving the complete nucleotide sequence of the HARS-1 fragment (see page 58). This sequence was determined shortly before the expiry of the priority year. Re-analysis of the experimental data has revealed that the sequence contained several errors.

A corrected sequence of the complete nucleotide sequence of the HARS-1 fragment is now provided.

It is requested that this correction of errors made by Applicants is allowed by the Patent Office in order to correct a part of the disclosure which is now known to be wrong.

Van der Toorren, Johannes Drs. European Patent Attorney

General Authorization No. 170

6

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Fig. 3 (amended)

DNA sequence of the autonomous-replicating sequence HARS1 from the methylotrophic yeast <u>Hansenula polymorpha</u>. The HARS1 represents a <u>SalI</u> fragment comprising 499 nucleotides. The dideoxy-sequencing method was employed.

CAACGCCGGA ATACGTCAGA AACAGCCGTG CCCCAGGGAC 80
CAGAAAGCCT ACTGGTGAGT ATGTTCTTTC GTGTGATTTT 120
TCCGAGGATG AGAACGACGA TAACGAGCAC AACTCGGAGT 160
CGGAGGACAC GCTTATTGCG TTGAACGCAG CCACATCAGC 200
AGGCTGTCAA GACTGAGTAT GGCCACAGAG CTGGATTCTC 240
GGCCTCATAC TCAAGACGTT AGTAAACTCC GTCTGCCAGA 280
AATTGCTGAC GAGGATGTAT AATAATAGAT GAATTACGAA 320
CAATTGTAGT TCAAAAAAAAT TTAGTAACAA TATTGTCTAG 360
ATGACAGATG TGCTGAAACC AGTGAACTCC AATAAACCAC 400
TCACCGCTAC CCAAGAGAAA CAGATCAGAG TGCTAGGGCC 440
TTGTTTCAGA GTACTACAAC GTTTACCAGA AGCTTGAGCA 480
AGTTCTCAAA CGCGGGTTTG (TCGAC)

500



(1) Publication number:

C12N13/53.

0 173 378

**A3** 

Office européen des brevets

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#### **EUROPEAN PATENT APPLICATION**

(21) Application number: 85201235.0

(22) Date of filing: 25.07.85

(51) Int. Ca.4: C 12 N 15/00

C 12 N 1/16, C 11 D 3/386 C 11 D 3/395, C 12 P 21/02 C 07 H 21/04, C 12 N 9/02

A request for correction of Fig. 3 has been filed pursuant to Rule 88 EPC. A decision on the request will be teken during the proceedings before the Examining Division.

- (30) Priority: 27.07.84 EP 84201114 07.02.85 GB 8503160
- (43) Date of publication of application: 05.03.86 Bulletin 86/10
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Use of oxidoreductases in bleaching and/or detergent compositions and their preperation by microorgenisms engineered by recombinent DNA technology.

(5) The structual genes and their regulatory DNA sequences of an alcohol oxidase (MOX) and a dihydroxyacetone synthase (DHAS) of Hansenula polymorpha have been isolated and the nucleotide sequences determined. The invention relates to the use of the MOX gene, as well as the use of the regulatory DNA sequences of MOX and/or DAS in combination with the MOX gene, optionally after modification thereof, or other oxidase genes, or other genes, to produce engineered microorganisms, in particular yeasts.

Said engineered microorganisms can produce oxidases or other enzymes in yields that allow industrial application on a large scale.

Moreover, said engineered microorganisms can produce oxidases having improved properties with respect to their application in oxidation reactions and/or in bleaching and detergent products.

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	DOCUMENTS CON	SIDERED TO BE RELEVAN	T	٦
Category	Citation of document v	with indication, where appropriate, evant passages	Relevant to claim	CLASSIFICATION OF THE APPLICATION (Int. CI.4)
х	EP-A- 0 103 887 * Claims 1-28 *	(AMGEN)	1	C 12 N 15/00 C 12 N 1/16 C 11 D 3/395
x		(NABISCO BRANDS)		C 11 D 3/386 C 12 P 21/02 C 07 H 21/04 C 12 N 9/02
	* Claims 1-7; pa		T	
۲	EP-A- 0 086 139	(TRANSGENE)		
	* Claims 1-38 *		1	
x	EP-A- 0 066 994	(ICI)		
	* Claims 1-17 *		1	
	February 1984, p 46308t; Columbus G. BRANLANT et a ning of the glyc phate dehydrogen lus stearothermo	eraldehyde-3-phos- ase genes of Bacil- philus and Escheri- heir expression in		TECHNICAL FIELDS SEARCHED (Int. CI 4)  C 12 N C 11 D
	* Abstract *		1	
	march 1982, page 63538j; Columbus C.A. LEE et al.: synthesis of enz	, Ohio, US "Plasmid-directed ymes required for port and utiliza-		
_	The present search report has b	een drawn up for all claims		
	Place of search The Hague	Date of completion of the search $17-04-1986$		Examiner DELANGHE
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	DOCUMENTS CONSIDI			lovas:	CLASSIFICATION OF THE
ategory	Citation of document with incontrol of relevant	dication, where appropriate, passages		elevant	APPLICATION (Int. Cl.4)
	<pre>% PROC. NATL. ACAD 78(12), 7336-40.  * Abstract *</pre>	. SCI. USA 1981	,   1		
		-			
x	CHEMICAL ABSTRACTS May 1981, page 375 Columbus, Ohio, US E. LOHMEIER et al. expression of the tase gene of Esche & CAN. J. BIOCHEM. 158-64.	: "Cloning and fumarate reductions coli."	1291		
	* Abstract *		1		
x	JOURNAL OF BIOCHER no.4, April 1982, Tokyo, JP M. IWAKURA et al. drofolate reducta richia coli K12."	pages 1205-121 : "Cloning of d se gene of Esch	ihy		TECHNICAL FIELDS SEARCHED (Int. CI.4)
	* Pages 1205-1211	* ·	1		
A	CHEMICAL ABSTRACT July 4, 1983, page 2206q; Columbus, M.J. WAITES et all acetone syntheses ketolase for form from the methylot Candida boidinii & MICROBIOL. 1983	Ohio, US  Ohio, US  a special tran  aldehyde fixati  crophic yeast  CBS 5777."	ns- Lon		
	* Abstract *		./.	1,35	
	The present search report has b	een drawn up tor all claims			
	Place of search	Date of completion of the	search		Examiner
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	OF UNITY OF INVENTION
	vision considers that the present European patent application does not comply with the requirement of unity of
	elates to severel invantions or groups of inventions,
	Claims 1-30: Process for preparing oxidoreductases by
,,	recombinant technology, their use, DNA sequences encoding oxidoreductases, process for preparing a transformed microorganism, microorganisms used
2)	Claims 35-39: DNA sequences coding for DHAS
	Claims 31-34,40-47: DNA sequences containing a regulon and a structural gene coding for a specific enzyme or other protein.  Process for preparing this enzyme.
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	report has been drawn up for those parts of the Europeen patent application which relets to the inventions in respect of which search teas here been paid.
	namely claims:
	None of the further accret fees has been paid within the fixed time limit. The present European asarch report has been drawn up for those parts of the European patent application which relate to the invention first mentioned in the claims.
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	DOCUMENTS CONSI	DERED TO BE R	ELEVANT		
alegory	Citation of document with of releva	indication, where approprint passages	rate,	Relevant to claim	CLASSIFICATION OF THE APPLICATION (Int. Cl.4)
	gene coding for m Hansenula polymor		lase in		
	* Whole document.	*		1-30	
1	_	· <b></b>			
?,X	NUCLEIC ACIDS RES 9, May 1985, page Oxford, GB Z.A. JANOWICZ et characterization encoding the majo latory enzyme fro phic yeast Hanser	al.: "Cloning of the DAS or methanol about the methy."	ng and gene assimi- lotro-		· ·
	* Whole document	*		1-39	
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			•		TECHNICAL FIELDS SEARCHED (Int. CI 4)
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	DOCUMENTS CON	SIDERED TO BE RELEVAN	τ	, <u>-</u>
Category		with indication, where appropriate, levant passages	Relevant to claim	CLASSIFICATION OF THE APPLICATION (Int. Cl.4)
А	March 29, 1982, 102478m; Columbon M. BRAVO et al. of methanol to pand hydrogen per & ADV. BIOTECHNO	: "Enzymic oxidation produce formaldehyde roxide." DL., (PROC. INT.		
	* Abstract *		1	
	August 3, 1981, 37855v; Columbus J. GEISSLER et a oxidases: An unu protein."	CTS, vol.95, no.5, page 291, ref.no.6, Ohio, US al.: "Yeast methanol asual type of flavo-81, 126(2), 152-6.		
	* Abstract *		1	TECHNICAL FIELDS SEARCHED (Int. CI 4)
?,X	WO-A- 84 04 539 SCHOOL DELFT)	(TECHNISCHE HOGE-		
	* Claims *	· 	1 ·	
x, x	WO-A- 85 01 063 SCHOOL DELFT)	(TECHNISCHE HOGE-		
	* Claims *		1	
	9, May 1985, pag Oxford, GB A.M. LEDEBOER et	SEARCH, vol.13, no. es 3063-3082; al.: "Molecular acterization of a		·
	The present search report has b	, 1	]	
	Place of search	Date of completion of the search	<del>                                     </del>	Examiner
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